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	(71) Applicant (for all designated States except US): THE FELLER UNIVERSITY [US/US]; 1230 York Aver York, NY 10021 (US).	Without international search rep upon receipt of that report.	oort and to be republished						
	(72) Inventors; and (75) Inventors/Applicants (for US only): MASURE, H (US/US); 430 East 63rd Street, Apartment 1/ York, NY 10021 (US). PEARCE, Barbara, J [AU/ East 63rd Street, Apartment 3N, New York, N (US). TUOMANEN, Elaine [US/US); 430 East 63 Apartment 12C, New York, NY 10021 (US).								

(54) Title: BACTERIAL EXPORTED PROTEINS AND ACELLULAR VACCINES BASED THEREON

#### (57) Abstract

The present invention relates to the identification of Gram positive bacterial exported proteins, and the genes encoding such proteins. In particular, the invention relates to adhesion associated exported proteins, and to antigens common to many or all strains of a species of Gram positive bacterium. The invention also relates to acultura vaccines to provide protection from Gram positive bacterium. The invention also relates to acultural vaccines to provide protection from Gram positive bacterium. The invention also because it is a species of grampers of the proteins, and to antibodies against such proteins for use in diagnosis and passive immune thermy). In specific embodiments, fragments of ten genes encoding exported proteins of S. preumoniae are disclosed, and the functional activity of some of these proteins in adherence is demonstrated.

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# BACTERIAL EXPORTED PROTEINS AND ACELLULAR VACCINES BASED THEREON

The research leading to the present invention was supported in part by the United

States Government, Grant No. R01-AI27913. The Government may have certain
rights in the invention.

#### CONTINUING INFORMATION

The present invention is a continuation-in-part of copending Application Serial No. 08/245,511, filed May 18, 1994, which is a continuation-in-part of copending Application Serial No. 08/116,541, filed September 1, 1993, each of which is incorporated by reference herein in its entirety, and applicants claim the benefit of the filing date of both applications pursuant to 35 U.S.C. § 120.

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### FIELD OF THE INVENTION

The present invention relates to the identification of bacterial exported proteins, and the genes encoding such proteins. The invention also relates to acellular vaccines to provide protection from bacterial infection using such proteins, and to antibodies against such proteins for use in diagnosis and passive immune therapy.

#### BACKGROUND OF THE INVENTION

25 Exported proteins in bacteria participate in many diverse and essential cell functions such as motility, signal transduction, macromolecular transport and assembly, and the acquisition of essential nutrients. For pathogenic bacteria, many exported proteins are virulence determinants that function as adhesins to colonize and thus infect the host or as toxins to protect the bacteria against the host's immune system (for a review, see Hoepelman and Tuomanen, 1992, Infect. Immun. 60:1729-33).

Since the development of the smallpox vaccine by Jenner in the 18th century,

vaccination has been an important armament in the arsenal against infectious microorganisms. Prior to the introduction of antibiotics, vaccination was the major hope for protecting populations against viral or bacterial infection. With the advent of antibiotics in the early 20th century, vaccination against bacterial infections became much less important. However, the recent insurgence of antibiotic-resistant strains of infectious bacteria has resulted in the reestablishment of the importance of anti-bacterial vaccines.

One possibility for an anti-bacterial vaccine is the use of killed or attenuated

bacteria. However, there are several disadvantages of whole bacterial vaccines,
including the possibility of a reversion of killed or attenuated bacteria to virulence
due to incomplete killing or attenuation and the inclusion of toxic components as
contaminants.

15 Another vaccine alternative is to immunize with the bacterial carbohydrate capsule. Presently, vaccines against Streptococcus pneumoniae employ conjugates composed of the capsules of the 23 most common serotypes of this bacterium, these vaccines are ineffective in individuals most susceptible to pathological infection — the young, the old, and the immune compromised — because of its inability to elicit a T cell immune response. A recent study has shown that this vaccine is only 50% protective for these individuals (Shapiro et al., 1991, N. Engl. J. Med. 325:1453-60).

An alternative to whole bacterial vaccines are acellular vaccines or subunit

25 vaccines in which the antigen includes a bacterial surface protein. These vaccines could potentially overcome the deficiencies of whole bacterial or capsule-based vaccines. Moreover, given the importance of exported proteins to bacterial virulence, these proteins are an important target for therapeutic intervention. Of particular importance are proteins that represent a common antigen of all strains of a particular species of bacteria for use in a vaccine that would protect against all strains of the bacteria. However, to date only a small number of exported proteins

of Gram positive bacteria have been identified, and none of these represent a common antigen for a particular species of bacteria.

A strategy for the genetic analysis of exported proteins in E. coli was suggested 5 following the description of translational fusions to a truncated gene for alkaline phosphatase (phoA) that lacked a functional signal sequence (Hoffman and Wright, 1985. Proc. Natl. Acad. Sci. U.S.A. 82:5107-5111). In this study, enzyme activity was readily detected in strains that had gene fusions between the coding regions of heterologous signal sequences and phoA indicating that translocation across the cytoplasmic membrane was required for enzyme activity. Subsequently, a modified transposon, TnphoA, was constructed to facilitate the rapid screening for translational gene fusions (Manoil and Beckwith, 1985, Proc. Natl. Acad. Sci. U.S.A. 82:8129-8133). This powerful tool has been modified and used in many Gram negative pathogens such as Escherichia coli (Guitierrez et al., 1987, J. Mol. Biol. 195;289-297), Vibrio cholera (Taylor et al., 1989, J. Bacteriol, 171:1870-1878), Bordetella pertussis (Finn et al., 1991, Infect Immun. 59:3273-9; Knapp and Mekalanos, 1988, J. Bacteriol, 170:5059-5066) and Legionella pneumophila (Albano et al., 1992, Mol. Microbiol, 6:1829-39), to yield a wealth of information from the identification and characterization of exported proteins. A similar strategy based on gene fusions to a truncated form of the gene for  $\beta$ -lactamase has been used to the same end (Broome-Smith et al., 1990, Mol. Microbiol. 4:1637-1644). A direct strategy for mapping the topology of exported proteins has also been developed based on "sandwich" gene fusions to phoA (Ehrmann et al., 1990, 87:7574-7578).

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For a variety of reasons, the use of gene fusions as a genetic screen for exported proteins in Gram positive organisms has met with limited success. Plasmid vectors that will create two or three part translational fusions to genes for alkaline phosphatase,  $\beta$ -lactamase and a-amylase have been designed for Bacillus subtilis and Lactococcus lacti (Payne and Jackson, 1991, J. Bacteriol. 173:2278-82; Perez et al., 1992, Mol. Gen. Genet, 234:401-11; Smith et al., 1987, J. Bacteriol.

169:3321-3328; Smith et al., 1988, Gene 70:351-361). Gene fusions between phoA and the gene for protein A (spa) from Staphylococcus aureus have been used to determine the cellular localization of this protein (Schneewind et al., 1992, Cell. 70:267-81). In that study, however, enzyme activity for alkaline phosphatase
was not reported.

Mutagenesis strategies in several streptococcal species have also been limited for several reasons. Efficient transposons similar to those that are the major tools to study Gram negative bacteria have not been developed for streptococcus. Insertion 10 duplication mutagenesis with non-replicating plasmid vectors has been a successful alternative for Streptococcus pneumoniae (Chen and Morrison, 1988, Gene. 64:155-164; Morrison et al., 1984, J. Bacteriol. 159:870). This strategy has led to the mutagenesis, isolation and cloning of several pneumococcal genes (Alloing et al., 1989, Gene. 76:363-8; Berry et al., 1992, Microb. Pathog. 12:87-93; Hui 15 and Morrison, 1991, J. Bacteriol. 173:372-81; Lacks and Greenberg, 1991, Gene. 104:11-7; Laible et al., 1989, Mol. Microbiol. 3:1337-48; Martin et al., 1992, J. Bacteriol, 174:4517-23; McDaniel et al., 1987, J. Exp. Med. 165:381-94; Prudhomme et al., 1989, J. Bacteriol, 171:5332-8; Prudhomme et al., 1991, J. Bacteriol. 173:7196-203; Puyet et al., 1989, J. Bacteriol. 171:2278-2286; Puyet et 20 al., 1990, J. Mol. Biol. 213:727-38; Radnis et al., 1990, J. Bacteriol, 172:3669-74; Sicard et al., 1992, J. Bacteriol. 174:2412-5; Stassi et al., 1981, Proc. Natl. Acad. Sci. U.S.A. 78:7028-7032; Tomasz et al., 1988, J. Bacteriol, 170:5931-5934; Yother et al., 1992, J. Bacteriol. 174:610-8).

25 Of note in the search for exported pneumococcal proteins that might be attractive targets for a vaccine is pneumococcal surface protein A (PspA) (see Yother et al., 1992, supra). PspA has been reported to be a candidate for a S. pneumoniae vaccine as it has been found in all pneumococci to date; the purified protein can be used to elicit protective immunity in mice; and antibodies against the protein confer passive immunity in mice (Talkington et al., 1992, Microb. Pathog. 13:343-355). However, PspA demonstrates antigenic variability between strains in

the N-terminal half of the protein, which contains the immunogenic and protection eliciting epitopes (Yother et al., 1992, supra). This protein does not represent a common antigen for all strains of S. pneumoniae, and therefore is not an optimal vaccine candidate.

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Recently, apparent fusion proteins containing PhoA were exported in species of Gram positive and Gram negative bacteria (Pearce and Masure, 1992, Abstr. Gen. Meet. Am. Soc. Microbiol. 92:127, abstract D-188). This abstract reports insertion of pneumococcal DNA upstream from the E. coli phoA gene lacking its signal sequence and promoter in a shuttle vector capable of expression in both E. coli and S. pneumoniae, and suggests that similar pathways for the translocation of exported proteins across the plasma membranes must be found for both species of bacteria.

- 15 Recent studies have shown that genetic transfer in several bacterial species relies on a signal response mechanism between individual cells. Conjugal plasmid transfer is mediated by homoserine lactones in Agrobacterium tumifaciens (Zhang et al., 1993, Scinece 362:446-448) and by small secreted polypeptides in Enterococcus faecalis (for a review, see Clewell, 1993, Cell 73:9-12). Low
- 20 molecular weight peptide activators have been described which induce transformation in S. pneumoniae (Tomasz, 1965, Nature 208:155-159; Tomasz, 1966, J. Bacteriol. 91:1050-61; Tomasz and Mosser, 1966, Proc. Natl. Acad. Sci. USA 55:58-66) and Streptococcus sanguis (Leonard and Cole, 1972, J. Bacteriol. 110:273-280; Pakula et al., 1962, Acta Microbiol. Pol. 11:205-222; Pakula and
- 25 Walczak, 1963, J. Gen. Microbiol. 31:125-133). A peptide activator which regulates both sporulation and transformation has been described for B. subtilis (Grossman and Losick, 1988, Proc. Natl. Acad. Sci. USA 85:4369-73). Furthermore, genetic evidence suggests that peptide permeases may be mediating these processes in both E. faecalis (Ruhfel et al., 1993, J. Bacteriol. 175:5253-59;
- 30 Tanimoto et al., 1993, J. Bacteriol. 175:5260-64) and B. subtilis (Rudner et al., 1991, J. Bacteriol. 173:1388-98).

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In S. pneumoniae, transformation occurs as a programmed event during a physiologically defined "competent" state. Induced by an unknown signal in a density dependent manner, cells exhibit a single wave of competence between 5 x 10° and 1-2 x 10° cfu / ml which is the beginning of logarithmic growth (Tomasz, 5 1966, supra). With induction, a unique set of competence associated proteins are expressed (Morrison and Baker, 1979, Nature 282:215-217) suggesting global regulation of transformation associated genes. Competent bacteria bind and transport exogenous DNA, which if homologous is incorporated by recombination into the genome of the recipient cell. Within one to two cell divisions, the 10 bacteria are no longer competent. As with induction, inactivation of competence occurs by an unknown mechanism.

The citation of references herein shall not be construed as an admission that such is prior art to the present invention.

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#### SUMMARY OF THE INVENTION

The present invention concerns genes encoding exported proteins in a Gram positive bacteria, and the proteins encoded by such genes. In particular, the invention provides for isolation of genes encoding Gram positive bacterial adhesion associated proteins, preferably adhesins, virulence determinants, toxins, or immunodominant proteins, and thus provides the genes and proteins encoded thereby. In another aspect, the exported protein can be an antigen common to many or all strains of a species of Gram positive bacteria, and that may be antigenically related to a homologous protein from a closely related species of bacteria. The invention also contemplates identification of proteins that are antigenically unique to a particular strain of bacteria. Preferably, the exported protein is an adhesin common to all strains of a species of Gram positive bacteria.

30 The invention further relates to a vaccine for protection of an animal subject from infection with a Gram positive bacterium comprising a vector containing a gene -7-

encoding an exported adhesion associated protein, or a gene encoding an exported protein which is an antigen common to many strains, of a species of a Gram positive bacterium operably associated with a promoter capable of directing of directing expression of the gene in the subject.

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In another aspect, the invention is directed to a vaccine for protection of an animal subject from infection with a Gram positive bacterium comprising an immunogenic amount of an exported adhesion associated protein, virulence determinant, toxin, or immunodominant protein of a Gram positive bacterium, or an immunogenic amount of an exported protein which is an antigen common to many strains of a species of Gram positive bacterium, and an adjuvant. Preferably, such a vaccine contains the protein conjugated covalently to a bacterial capsule or capsules from one or more strains of bacteria. More preferably, the capsules from all the common strains of a species of bacteria are included in the vaccine.

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Alternatively, the protein can be used to immunize an appropriate animal to generate polyclonal or monoclonal antibodies, as described in detail below. Thus, the invention further relates to antibodies reactive with exported proteins of Gram positive bacteria. Such antibodies can be used in immunoassays to diagnose 20 infection with a particular strain or species of bacteria. Thus, strain-specific exported proteins can be used to generate strain-specific antibodies for diagnosis of infection with that strain. Alternatively, common antigens can be used to prepare antibodies for the diagnosis of infection with that species of bacterium. In a specific aspect, the species of bacterium is S. pneumoniae. The antibodies can also be used for passive immunization to treat an infection with Gram positive bacteria

Thus, it is an object of the present invention to provide genes encoding exported proteins of Gram positive bacteria. Preferably, such genes encode adhesion

30 associated proteins, virulence determinants, toxins, or immunodominant proteins that are immunogenic. Preferably, the protein is an antigen common to many

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strains of a species of Gram positive bacterium, as the products of such genes are particularly attractive vaccine candidates.

It is a further object of the invention to provide an acellular vaccine against a

Gram positive bacterium, thus overcoming the deficiencies of whole killed or
attenuated bacterial vaccines and capsular vaccines.

Another object of the present invention is to provide a capsular vaccine that elicits a helper T cell immune response.

It is yet a further object of the invention to provide for the diagnosis of infection with a Gram positive bacterium.

Another object of the invention is to provide for passive immune therapy for a

15 Gram positive bacterial infection, particularly for an infection by an antibiotic
resistant bacterium.

#### BRIEF DESCRIPTION OF THE DRAWINGS

20 FIGURE 1. Construction of PhoA fusion vectors designed for the mutation and genetic identification of exported proteins in S. pneumoniae. (A) The 2.6 kB fragment of pPHO7 containing a truncated form of phoA was inserted into either the Smal or BamHl sites of pJDC9 to generate pHRM100 and pHRM104 respectively. T1T2 are transcription terminators and the arrows indicate gene orientation. (B) Mechanism of insertion duplication mutagenesis coupled to gene fusion. PhoA activity depends on the cloning of an internal gene fragment that is in-frame and downstream from a gene that encodes an exported protein. Transformation into S. pneumoniae results in duplication of the target fragment and subsequent gene disruption.

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FIGURE 2. Detection and trypsin susceptibility of PhoA fusions in S.

pneumoniae. Total cells lysates (50 µg of protein) from R6x (lane 1; parental strain): SPRU98 (lane 2); SPRU97 (lane 3); and SPRU96 (lane 4) were applied to an 8-25% SDS polyacrylamide gel. Proteins were transferred to nitrocellulose membranes and probed with anti-PhoA antibody. Antigen-antibody complexes

- 5 were detected by enhanced chemiluminescence with an appropriate peroxidase conjugated second antibody. SPRU96 and 97 contain the plasmids pHRM100 and pHRM104 randomly integrated in the chromosome. Molecular weight standards are indicated on the left. Whole bacteria from strain SPRU98 were treated with (lane 5) and without (lane 6) 50 μg / ml of trypsin for 10 min. at 37 °C. Both
- 10 samples were treated with a 40 fold molar excess of soy bean trypsin inhibitor. The total cell lysates (50 µg protein) were probed for immunoreactive material to PhoA as described above. Molecular weight standards are indicated on the left.
- FIGURE 3. PhoA fusion products are more stable when bacteria are grown in the presence of disulfide oxidants. Cultures of SPRU98 were grown in the presence of either 600 µM 2-hydroxyethel disulfide (lane 1), 10 µM DsbA (lane 2) or without any additions (lane 3). Total cell lysates (50 µg of protein) were applied to an 8 25% SDS polyacrylamide gel. The proteins were then probed for immunoreactive material with anti PhoA antibody as described in Figure 2.

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- FIGURE 4. Derived amino acid sequences for the genetic loci recovered from PhoA<sup>+</sup> pneumococcal mutants. Each of the plasmids recovered from the nine PhoA<sup>+</sup> strains of S. pneumoniae (see Table 1) were transformed into E. coli and had 400 to 700 base pair inserts. Using a primer to the 5' end of phoA,
- 25 approximately 200 to 500 base pairs of pneumococcal DNA immediately upstream of phoA was sequenced from each plasmid and an in-frame coding region with PhoA was established. The derived amino acid sequences from the fusions are presented for Exp1 [SEQ ID NO:2], Exp2 [SEQ ID NO:24], Exp3 [SEQ ID NO:6], Exp4 [SEQ ID NO:8], Exp5 [SEQ ID NO:10], Exp6 [SEQ ID NO:12],
- 30 Exp7 [SEQ ID NO:14], Exp8 [SEQ ID NO:16], and Exp9a [SEQ ID NO:18].
  The derived sequence from the 5' end of the insert from Exp9 is also presented in

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Exp9b [SEQ ID NO:20].

FIGURE 5. Sequence alignments of the derived amino acid sequences from the Exp loci recovered from PhoA+ mutants. The highest scoring match for each 5 insert is presented. The percent identity (%ID) and percent similarity (%SIM) for each alignment is presented on the right. (A) Expl ISEO ID NO:21 and AmiA from S. pneumoniae [SEO ID NO:23] (Alloing et al., 1990, Mol. Microbiol. 4:633-44). B) Exp2 [SEQ ID NO:24] and PonA from S. pneumoniae [SEO ID NO:241 (Martin et al., 1992, J. Bacteriol, 174:4517-23). C) Exp3 [SEO ID 10 NO:25] and PilB from N. gonorrhoeae [SEQ ID NO:26] (Taha et al., 1988, EMBO J. 7:4367-4378). The conserved histidine (Hans) in PilB is not present in Exp3 but is replaced by asparagine (N124). D) Exp4 [SEQ ID NO:27] and CD4B from tomato [SEQ ID NO:28] (Gottesman et al., 1990, Proc. Natl. Acad. Sci. U.S.A. 87:3513-7). E) Exp5 [SEO ID NO:29] and PtsG from B. subtilis [SEO 15 ID NO:30] (Gonzy-Tréboul et al., 1991, Mol. Microbiol. 5:1241-1294). F) Exp6 [SEQ ID NO:31] and GlpD from B. subtilis [SEQ ID NO:32] (Holmberg et al., 1990, J. Gen. Microbiol, 136-2367-2375), G) Exp7 [SEO ID NO:33] and MgtB from S. typhimurium [SEQ ID NO:34] (Snavely et al., 1991, J. Biol. Chem. 266:815-823). The conserved aspartic acid (D<sub>554</sub>) required for autophosphorylation 20 is also present in Exp7 (D<sub>37</sub>). H) Exp8 [SEO ID NO:35] and CyaB from B. pertussis [SEQ ID NO:36] (Glaser et al., 1988, Mol. Microbiol. 2:1930; Glaser et al., 1988, EMBO J. 7:3997-4004). I) Exp9 and DeaD from E. coli (Toone et al., 1991, J. Bacteriol, 173;3291-3302). The top sequence from Exp9 [SEO ID NO:37] is derived from the 5' end of the recovered plasmid insert, and compared 25 to DeaD 135-220 [SEO ID NO:38]. The bottom sequence from Exp9 [SEO ID NO:201 is derived from the 3' end of the recovered plasmid insert just unstream from phoA, and is compared with DeaD 265-342 [SEQ ID NO:39]. The

30 FIGURE 6. Subcellular localization of the Exp9-PhoA fusion. The membrane (lane 1) and cytoplasmic (lane 2) fractions (50 µg of protein for each sample) of

conserved DEAD sequence is highlighted.

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SPRU17 were applied to a 10-15% SDS polyacrylamide gel. The proteins were transferred to nitrocellulose and probed with anti-PhoA antibody. Molecular weight standards are indicated on the left.

- 5 FIGURE 7. Adherence of type 2 AII ( ) or unencapsulated R6 (O) pneumococci to alveolar Type II cells of rabbit. The adherence assay was performed as described in Example 2, infra.
- FIGURE 8. Titration of the adherence of pneumococcal mutants to human umbilical vein endothelial cells (HUVEC). The mutant strains tested are listed on Table 1. Mutation of exp1, strain SPRU98 (●); exp2, strain SPRU64 (○); exp3, strain SPRU40 (■); exp10, strain SPRU25 (⑤); and amiA, strain SPRU121 (◆) resulted in a decrease in the ability of the mutant strain to adhere. Strain R6 (■) is wildtype S. pneumoniae.

FIGURE 9. Adherence of pneumococcal mutants to lung Type II cells. The exported gene mutation and strain designations are as described for Figure 8.

FIGURE 10. Nucleotide and deduced amino acid sequences for the genetic locus
recovered from the SPRU25 mutant, exp10. The nucleotide sequence was
obtained as described in Figure 4 and in Example 1, infra.

FIGURE 11. Nucleotide (SEQ ID NO: 46) and derived protein (SEQ ID NO: 47) sequences of plpA. The lipoprotein modification consensus sequence is underlined with an asterisk above the cysteine residue where cleavage would occur.

Downstream from the coding region a potential rho independent transcription terminator is underlined. The positions of the PhoA fusions at Leu<sub>197</sub> in SPRU58 and Asp<sub>492</sub> in SPRU98 are indicated. (Genbank accession number: TO BE ASSIGNED).

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FIGURE 12. Sequence analysis of peptide binding proteins. A; Sequence

alignment of PlpA (SEQ ID NO:47) and AmiA (SEQ ID NO:48). Identical residues are boxed. B; Sequence alignments for the substrate binding proteins from the permeases of different bacterial species: PlpA, S. pneumoniae (this study); AmiA, S. pneumoniae. The reported sequence for amiA (Alloing et al.,

- 5 1990, Mol. Microbiol. 4:633-644) has now been changed due to a sequencing error and the corrected sequence is now in Genbank); Spo0KA, B. subtilis (Perego et al., 1991, Mol. Microbiol. 5:173-185; Rudner et al., 1991, J. Bacteriol. 173:1388-98); HbpA, H. influenzae (Hanson et al., 1992, Infect. Immun. 60:2257-66); DciAE, B. subtilis (Mathiopoulos et al., 1991, Mol. Microbiol. 5:1903-13);
- OppA (Ec), E. coli (Kashiwagi et al., 1990, J. Biol. Chem. 265:8387-91); TraC, E. faecalis (Tanimoto et al., 1993, J. Bacteriol. 175:5260-64); DppA, E. coli (Abouhamad et al., 1991, Mol. Microbiol. 5:1035-47); PrgZ, E. faecalis (Ruhfel et al., 1993, J. Bacteriol. 175:5253-59); OppA (St) S. typhimurium (Hiles et al., 1987, J. Mol. Biol. 195:125-142) and SarA, S. gordonii. The derived amino acid sequences were aligned with the MACAW software package (Schuler et al., 1993,
  - Proteins Struct. Funct. Genet. 9:180-190). The black boxes and hatched boxes denote regions of high sequence similarity with probability values less than or equal to 1.3 x 10<sup>-7</sup>, with the effective size of the space searched derived from the lengths of all the sequences in the database.

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FIGURE 13. Subcellular localization and labeling of PlpA-PhoA. Upper panel: Subcellular fractions (50 μg of total protein) from SPRU98 (PhoA<sup>+</sup>, pHRM104::*plpA*) were applied to an 8-25% SDS polyacrylamide gel, transferred to a nitrocellulose membrane and probed with anti-PhoA antisera. Bound

- 25 antibodies were detected with a peroxidase conjugated second antibody and visualized with enhanced chemiluminescence. Lanes are A, culture supernatant; B, membranes; C, cytoplasm; and D, cell wall. Lower panel: Anti-PhoA immunoprecipitates of total cell lysates from bacteria grown in a chemically defined media with PH palmitic acid were applied to an 8-25% SDS
- 30 polyacrylamide gel, transferred to a nitrocellulose membrane and subjected to autoradiography. Lanes are E, parental strain R6x; F, SPRU100 (PhoA\*,

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pHRM104::zzz); and G, SPRU98 (PhoA+, pHRM104::plpA). The arrow marks the 93 kDa band that corresponds to the immunoprecipitated PlpA-PhoA fusion protein.

- 5 FIGURE 14. Northern analysis of pneumococcal peptide permases. RNA (10 μg) prepared from SPRU107 (pJDC9::plpA) (lanes A and C) and R6x (lanes B and D) was hybridized to DNA probes from plpA (lanes A and B) or amiA (lanes C and D). Molecular weights are indicated.
- 10 FIGURE 15. Transformation efficiency of pneumococcal permease mutants. Various strains containing the depicted chromosomal gene constructs with lesions in either plpA or ami were assayed for the incorporation of a chromosomal streptomycin resistance marker as a measure of transformation efficiency. Transformation efficiency of each strain is presented as a percent of the parental strain, R6x, which routinely produces 0.3% Str transformants in the total population of transformable cells. Values presented are the average of at least three data points with the standard error of the mean. The results are representative of assays performed on three separate occasions. E is erythromycin resistance encoded by the vector.

FIGURE 16. Competence profiles of pneumococcal permease mutants. The percentage of transformable cells was determined at specific ODs during early logarithmic growth for R6x n, SPRU1071 (pJDC9::plpA), and SPRU114 s (pJDC9::amiA). The results are representative of three separate experiments.

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FIGURE 17. Effect of a mutation in plpA on the expression of the competence regulated rec locus. Alkaline phosphatase activity was measured for SPRU100, n (PhoA+, pHRM104::exp10) and SPRU156, s (PhoA+, pHRM104::exp10; pWG5::plpA) during logarithmic growth of pneumococcus which produces a normal competence cycle. Each value is the average of two data points with a standard error of the mean that did not exceed 10% of that point. These results are

representative of three independent experiments.

FIGURE 18. Physical map of plpA and recombinant plasmids generated from various cloning procedures. Plasmids with the preface pH contain inserts in the 5 PhoA vector pHRM104 while plasmids with the preface pJ contain inserts in the vector pIDC9. Most plasmids were created by "chromosome walking" with the integrated plasmid pJplp1. The plasmid pJplp9 was created by "homology cloning" with the oligonucleotides lipo1 and P1. See experimental procedures for details. Restriction endonuclease sites are shown: H (HindIII), Hc (HincII), E 10 (EcoRI), K (KpnI), P (PstI), R (EcoRV), Sau (SauIIIa), S (SphI).

FIGURE 19. Adherence of R6 wild-type ( $\square$ ) and Pad1 mutant ( $\blacksquare$ ) pneumococci to type II lung cells. This assay was performed as described in Example 2.

FIGURE 20. (A) Subcellular localization of Pad1-PhoA fusion detected by Western analysis with anti-PhoA antisera. The cells were separated into the membrane components (Lanes A-C) and cytoplasmic components (Lanes D-F).
 Lanes A,D - R6 wild-type (parent) cells; B,E - Pad1 mutant cells; C,F -- Pad1b mutant cells. (B) Probe of bacterial lysate with antibody to whole bacteria by
 Western analysis. Lanes A, B and C correspond to (A). The Pad1 mutants lack a 17 kDa immunogenic membrane associated protein found in the R6 bacteria.

FIGURE 21. Adherence of R6 bacteria and Pad1 mutants grown in the presence and absence of acetate. Growth in acetate corrects the Pad1 adherence defect.

FIGURE 22. Growth of the Padl mutant and R6 bacteria in the presence or absence of acetate. The Padl mutant was grown in chhemically defined growth medium for S. pneumodiae in the presence of 0% ( $\bigcirc$ ), 0.1% ( $\bigcirc$ ) and 0.5% ( $\square$ ) acetate. R6 was grown in the presence of 0% (square plus) and 0.5% ( $\triangle$ ).

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25

FIGURE 23. Nucleotide (SEQ ID NO:55) and deduced amino acid sequences of

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Pad1 (SEQ ID NO:56); also termed poxB. The putative ribosome binding site, -10, and -35 sites are underlined, and the start codon is labeled.

#### DETAILED DESCRIPTION OF THE INVENTION

5

In accordance with the present invention there may be employed conventional molecular biology, microbiology, and recombinant DNA techniques within the skill of the art. Such techniques are explained fully in the literature. See, e.g., Sambrook, Fritsch & Maniatis, "Molecular Cloning: A Laboratory Manual,"

- 10 Second Edition (1989) Cold Spring Harbor Laboratory Press, Cold Spring Harbor, New York (herein "Sambrook et al., 1989"); "DNA Cloning: A Practical Approach," Volumes I and II (D.N. Glover ed. 1985); "Oligonucleotide Synthesis" (M.J. Gait ed. 1984); "Nucleic Acid Hybridization" [B.D. Hames & S.J. Higgins eds. (1985)]; "Transcription And Translation" [B.D. Hames & S.J.
- 15 Higgins, eds. (1984)]; "Animal Cell Culture" [R.I. Freshney, ed. (1986)]; "Immobilized Cells And Enzymes" [IRL Press, (1986)]; B. Perbal, "A Practical Guide To Molecular Cloning" (1984).

Therefore, if appearing herein, the following terms shall have the definitions set 20 out below.

A "replicon" is any genetic element (e.g., plasmid, chromosome, virus) that functions as an autonomous unit of DNA replication in vivo, i.e., capable of replication under its own control.

25

A "vector" is a replicon, such as plasmid, phage or cosmid, to which another DNA segment may be attached so as to bring about the replication of the attached segment.

30 The term " viral vector" refers to a virus containing a recombinant nucleic acid, whereby the virus can introduce the recombinant nucleic acid to a cell, i.e., the

virus can transform the cell. According to the present invention, such vectors may have use for the delivery of a nucleic acid-based vaccine, as described herein.

A cell has been "transformed" by exogenous or heterologous DNA when such

5 DNA has been introduced inside the cell. The transforming DNA may or may not
be integrated (covalently linked) into chromosomal DNA making up the genome of
the cell. In prokaryotes, yeast, and mammalian cells for example, the
transforming DNA may be maintained on an episomal element such as a plasmid.
A "clone" is a population of cells derived from a single cell or common ancestor

10 by mitosis.

A "nucleic acid molecule" refers to the phosphate ester polymeric form of ribonucleosides (adenosine, guanosine, uridine or cytidine; "RNA molecules") or deoxyribonucleosides (deoxyadenosine, deoxyguanosine, deoxythymidine, or deoxycytidine; "DNA molecules") in either single stranded form, or a double-stranded helix. Double stranded DNA-DNA, DNA-RNA and RNA-RNA helices are possible. The term nucleic acid molecule, and in particular DNA or RNA molecule, refers only to the primary and secondary structure of the molecule, and does not limit it to any particular tertiary forms. Thus, this term includes double-stranded DNA found, inter alia, in linear or circular DNA molecules (e.g., restriction fragments), viruses, plasmids, and chromosomes. In discussing the structure of particular double-stranded DNA molecules, sequences may be described herein according to the normal convention of giving only the sequence in the 5' to 3' direction along the nontranscribed strand of DNA (i.e., the strand 25 having a sequence homologous to the mRNA). A "recombinant DNA molecule" is a DNA molecule that has undersone a molecular biological manipulation.

A nucleic acid molecule is "hybridizable" to another nucleic acid molecule, such as a cDNA, genomic DNA, or RNA, when a single stranded form of the nucleic acid molecule can anneal to the other nucleic acid molecule under the appropriate conditions of temperature and solution ionic strength (see Sambrook et al., 1989.

supra). The conditions of temperature and ionic strength determine the "stringency" of the hybridization. Hybridization requires that the two nucleic acids contain complementary sequences, although depending on the stringency of the hybridization, mismatches between bases are possible. The appropriate
stringency for hybridizing nucleic acids depends on the length of the nucleic acids and the degree of complementation, variables well known in the art. Preferably a minimum length for a hybridizable nucleic acid is at least about 10 nucleotides; more preferably at least about 15 nucleotides.

10 A DNA "coding sequence" is a double-stranded DNA sequence which is transcribed and translated into a polypeptide in vivo when placed under the control of appropriate regulatory sequences. The boundaries of the coding sequence are determined by a start codon at the 5' (amino) terminus and a translation stop codon at the 3' (carboxyl) terminus. A coding sequence can include, but is not limited to, prokaryotic sequences, cDNA from eukaryotic mRNA, genomic DNA sequences from eukaryotic (e.g., mammalian) DNA, and even synthetic DNA sequences. If the coding sequence is intended for expression in a eukaryotic cell, a polyadenylation signal and transcription termination sequence will usually be located 3' to the coding sequence.

20

Transcriptional and translational control sequences are DNA regulatory sequences, such as promoters, enhancers, terminators, and the like, that provide for the expression of a coding sequence in a host cell. In eukaryotic cells, polyadenylation signals are control sequences.

25

A "promoter sequence" is a DNA regulatory region capable of binding RNA polymerase in a cell and initiating transcription of a downstream (3' direction) coding sequence. For purposes of defining the present invention, the promoter sequence is bounded at its 3' terminus by the transcription initiation site and extends upstream (5' direction) to include the minimum number of bases or elements necessary to initiate transcription at levels detectable above background,

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Within the promoter sequence will be found a transcription initiation site (conveniently defined for example, by mapping with nuclease S1), as well as protein binding domains (consensus sequences) responsible for the binding of RNA polymerase. Eukaryotic promoters will often, but not always, contain "TATA" boxes and "CAT" boxes.

A coding sequence is "under the control" of transcriptional and translational control sequences in a cell when RNA polymerase transcribes the coding sequence into mRNA, which is then translated into the protein encoded by the coding 10 sequence.

A "signal sequence" can be included before the coding sequence. This sequence encodes a signal peptide, N-terminal to the polypeptide, that directs the host cell to translocate the polypeptide to the cell surface or secrete the polypeptide into the media, and this signal peptide is selectively degraded by the cell upon exportation. Signal sequences can be found associated with a variety of proteins native to prokarvotes and eukarvotes.

As used herein, the term "exported protein" refers to a protein that contains a signal sequence, and thus is found associated with or outside of the cell membrane. Thus, secreted proteins, integral membrane proteins, surface proteins, and the like fall into the class of exported proteins. The term "surface protein" as used herein is specifically intended to refer to a protein that is accessible at the cell surface, e.g., for binding with an antibody.

25

An "adhesion associated protein" is a protein that is directly or indirectly involved in adherence of bacteria to target cells, such as endothelial cells or lung cells. The term "adhesion associated protein" includes proteins that may have other functional activities, such as motility, signal transduction, cell wall assembly, or

macromolecular transport. An "adhesin" is an adhesion-associated protein found on the surface of a cell, such as a bacterium, that is directly involved in adherence, and thus effects some degree of adherence or adhesion to another cell.

Of particular importance to the present invention are adhesins of Gram positive bacteria that promote adhesion to eukaryotic cells, i.e., that are involved in bacterial virulence. Adhesins, in order to be effective in promoting adherence, should be surface proteins, i.e., be accessible at the surface of the cell.

Accessibility is also important to determine antigenicity. A vaccine that elicits antibodies against an adhesin can provide antibodies that bind to an accessible antigenic determinant and directly interfere with adherence, thus preventing infection. An adhesin of the invention need not be the only adhesin or adhesion mediator of a Gram positive bacteria, and the term contemplates any protein that demonstrates some degree of adhesion activity, whether relatively strong or relatively weak.

A "virulence determinant" is any bacterial product required for bacterial survival

15 within an infected host. Thus, virulence determinants are also attractive vaccine

candidates since neutralization of a virulence determinant can reduce the virulence

of the bacteria.

A "toxin" is any bacterial product that actively damages an infected host. Thus,

20 bacterial toxins are important targets for an immune response in order to neutralize
their toxicity.

A molecule is "antigenic" when it is capable of specifically interacting with an antigen recognition molecule of the immune system, such as an immunoglobulin (antibody) or T cell antigen receptor. An antigenic polypeptide contains at least about 5, and preferably at least about 10, amino acids. An antigenic portion of a molecule can be that portion that is immunodominant for antibody or T cell receptor recognition, or it can be a portion used to generate an antibody to the molecule by conjugating the antigenic portion to a carrier molecule for immunization. A molecule that is antigenic need not be itself immunogenic, i.e., capable of eliciting an immune response without a carrier.

- 20 -

A composition comprising "A" (where "A" is a single protein, DNA molecule, vector, etc.) is substantially free of "B" (where "B" comprises one or more contaminating proteins, DNA molecules, vectors, etc.) when at least about 75% by weight of the proteins, DNA, vectors (depending on the category of species to which A and B belong) in the composition is "A". Preferably, "A" comprises at least about 90% by weight of the A+B species in the composition, most preferably at least about 99% by weight. It is also preferred that a composition, which is substantially free of contamination, contain only a single molecular weight species having the activity or characteristic of the species of interest.

10

The phrase "pharmaceutically acceptable" refers to molecular entities and compositions that are physiologically tolerable and do not typically produce an allergic or similar untoward reaction, such as gastric upset, dizziness and the like, when administered to a human. Preferably, as used herein, the term

15 "pharmaceutically acceptable" means approved by a regulatory agency of the Federal or a state government or listed in the U.S. Pharmacopeia or other generally recognized pharmacopeia for use in animals, and more particularly in humans. The term "carrier" refers to a diluent, adjuvant, excipient, or vehicle with which the compound is administered. Such pharmaceutical carriers can be sterile liquids, such as water and oils, including those of petroleum, animal, vegetable or synthetic origin, such as peanut oil, soybean oil, mineral oil, sesame oil and the like. Water or aqueous solution saline solutions and aqueous dextrose and glycerol solutions are preferably employed as carriers, particularly for

25

injectable solutions.

The term "adjuvant" refers to a compound or mixture that enhances the immune response to an antigen. An adjuvant can serve as a tissue depot that slowly releases the antigen and also as a lymphoid system activator that non-specifically enhances the immune response (Hood et al., Immunology, Second Ed., 1984, 30 Benjamin/Cummings: Menlo Park, California, p. 384). Often, a primary challenge with an antigen alone, in the absence of an adjuvant, will fail to elicit a

humoral or cellular immune response. Adjuvants include, but are not limited to, complete Freund's adjuvant, incomplete Freund's adjuvant, saponin, mineral gels such as aluminum hydroxide, surface active substances such as lysolecithin, pluronic polyols, polyanions, peptides, oil or hydrocarbon emulsions, keyhole limpet hemocyanins, dinitrophenol, and potentially useful human adjuvants such as BCG (bacille Calmette-Guerin) and Corynebacterium parvum. Preferably, the adjuvant is pharmaceutically acceptable.

In its primary aspect, the present invention concerns the identification and isolation of a gene encoding an exported protein in a Gram positive bacteria. The exported protein can be a protein of unknown or of known function. Herein, all such exported proteins, whether of known or of unknown function, are referred to as "Exp" (for exported protein), and the genes encoding such proteins are referred to as "exp" genes. In particular, the invention provides for isolation of genes encoding Gram positive bacterial adhesion associated proteins, preferably adhesins, virulence determinants, toxins and immunodominant antigens. Preferably, the exported protein can be an antigen common to all strains of a species of Gram positive bacteria, or that may be antigenically related to a homologous protein from a closely related species of bacteria. The invention also contemplates identification of proteins that are antigenically unique to a particular strain of bacteria. Preferably, the exported protein is an adhesin common to all strains of a species of Gram positive bacteria, in particular, S. pneumoniae.

In particular, the invention concerns various exported proteins of *S. pneumoniae*25 (see Table 1, infra), some of which demonstrate activity as adhesins. In specific embodiments, the invention provides gene fragments of the following exported proteins: Exp1 [SEQ ID NO:2], the full length sequence of which, termed Plp1 [SEQ ID NO:47], is also provided, encoded by exp1 [SEQ ID NO:1] and plp1 [SEQ ID NO:46], respectively, a protein that appears to be related to the permease family of proteins and which is therefore surprisingly associated with adhesion; Exp2 [SEQ ID NO:3], encoded by exp2 [SEQ ID NO:4], which nucleic acid

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sequence is identical to ponA, which encodes penicillin-binding protein 1A (Martin et al., 1992, J. Bacteriol, 174:4517-4523), and which is unexpectedly associated with adhesion; Exp3 [SEQ ID NO:6], encoded by exp3 [SEQ ID NO:5], which is associated with adhesion; Exp4 (SEQ ID NO:8), encoded by exp4 [SEQ ID 5 NO:7), which is associated with adhesion; Exp5 [SEQ ID NO:10], encoded by exp5 [SEO ID NO:9]; Exp6 [SEQ ID NO:12], encoded by exp6 [SEO ID NO:11]; Exp7 [SEO ID NO:14], encoded by exp7 [SEO ID NO:13]; Exp 8 [SEO ID NO:161, encoded by exp8 [SEO ID NO:15]; Exp9 [SEO ID NOS, 18 and 20]. encoded by exp9 [SEO ID NOS. 17 and 19, respectively]; Exp10 [SEO ID 10 NO:221, encoded by exp10 [SEO ID NO:21]; and Pad1 [SEO ID NO:56], encoded by pad1 [SEQ ID NO:55], which is a pyruvate oxidase homolog. The strain designations of mutant bacteria in which the Exp1-9 proteins were identified are disclosed in Table 1. The strain designation of the mutant in which Exp10 was identified is SPRU25. Applicants have also isolated a mutant S. pneumoniae 15 (SPRU121) in which the amiA gene encoding the AmiA protein has been mutated. and have demonstrated for the first time that this is an adhesion associated protein, and thus, that this protein can be used in a vaccine to elicit an anti-adhesionassociated protein immune response.

20 Once the genes encoding exported proteins are isolated, they can be used directly as an in vivo nucleic acid-based vaccine. Alternatively, the nucleotide sequence of the genes can be used to prepare oligonucleotide probes or primers for polymerase chain reaction (PCR) for diagnosis of infection with a particular strain or species of Gram positive bacterium.

25

Alteratively, the proteins encoded by the isolated genes can be expressed and used to prepare vaccines for protection against the strain of bacteria from which the exported protein was obtained. If the exported protein is an adhesion associated protein, such as an adhesin, it is a particularly attractive vaccine candidate since immunity can interfere with the bacterium's ability to adhere to host cells, and thus infect, i.e., colonize and survive, within host organism. If the exported

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protein is a virulence determinant, immunity can interfere with virulence. If the exported protein is a toxin, immunity can interfere with toxicity. More preferably, the exported protein is an antigen common to all or almost all strains of a particular species of bacterium, and thus is an ideal candidate for a vaccine

5 against all or almost all strains of that species. In a specific embodiment, the species of bacterium is S. pneumoniae.

Alternatively, the protein can be used to immunize an appropriate animal to generate polyclonal or monoclonal antibodies, as described in detail below. Such antibodies can be used in immunoassays to diagnose infection with a particular strain or species of bacteria. Thus, strain-specific exported proteins can be used to generate strain-specific antibodies for diagnosis of infection with that strain. Alternatively, common antigens can be used to prepare antibodies for the diagnosis of infection with that species of bacterium. In a specific aspect, the species of 15 bacterium is S. pneumoniae.

In yet another embodiment, if the Exp is an adhesin, the soluble protein can be administered to a subject suspected of suffering an infection to inhibit adherence of the hacterium.

20

#### Isolation of Genes for Exported Proteins

The present invention provides a number of gene fragments that can be used to obtain the full length gene encoding exported Gram positive bacterial antigens, in 25 particular exported adhesins.

The invention further provides a method, using a vector that encodes an indicator protein that is functional only when exported from a bacterium, such as the *phoA* vector described herein, to screen for genes encoding exported pneumococcal proteins. For example, a truncated form of *phoA* can be placed in a pneumococcal shuttle vector, such as vector pJDC9 (Chen and Morrison, 1988, Gene 64:155-

164). A cloning site containing a unique restriction site, e.g., Smal or BamHl can be located immediately 5' to phoA, to allow insertion of DNA that may encode an export protein. Preferably, the cloning sites in the vector are flanked by two restriction sites to facilitate easy identification of an insert. In a specific
5 embodiment, the restriction site is a Kpnl site, although any restriction endonuclease can be used. Gene fragments encoding Exp's are selected on the basis of blue staining around the bacterium, which is indicative of export of the PhoA enzyme. The exp-phoA fusion genes can be expressed in E. coli, although a promoter fusion may be required in this instance. When integrated into the
10 genome of a Gram positive organism, the exp-phoA fusion gene is a translational fusion involving duplication mutagenesis, and expressed in a Gram positive bacterium. In a specific embodiment, pneumococcal export proteins are identified with this technique, which requires cloning of an internal gene fragment within the vector prior to integration.

15

In a further embodiment, screening for genes encoding exported adhesion associated proteins can be performed on PhoA-positive transformants by testing for loss of adherence of a Gram positive bacterium to a primary cell or a cell line to which it normally adheres. Such adhesion assays can be performed on any 20 eukaryotic cell line. Preferably, if infection of humans is important, the cell or cell line is derived from a human source or has been demonstrated to behave like human cells in a particular in vitro assay. Suitable cells and cell lines include, but are not limited to, endothelial cells, lung cells, leukocytes, buccal cells, adenoid cells, skin cells, conjunctivial cells, ciliated cells, and other cells representative of 25 infected organs. As demonstrated in an example, infra, a human umbilical vein endothelial cell (HUVEC) line, which is available from Clonetics (San Diego. CA), can be used. In another example, infra, lung Type II alveolar cells, which can be prepared as described in Example 2 or can be obtained as a cell line available from the American Type Culture Collection (ATCC) under accession 30 number ATCC A549, are used. Alternatively, adherence to human monocytederived macrophages, obtained from blood, can be tested. Other target cells,

especially for *S. pneumoniae*, are oropharyngeal cells, such as buccal epithelial cells (Andersson et al. (1988, Microb. Pathogen. 4:267-278; 1983, J. Exp. Med. 158:559-570; 1981, Infect. Immun. 32:311-317).

5 Generally, any adherence assay known in the art can be used to demonstrate loss of adhesion due to mutagenesis of the Exp. One such assay follows: The cells to which adherence is to be assayed are cultured for 4-8 days (Wright AND Silverstein, 1982, J. Exp. Med. 156:1149-1164) and then transferred to Terasaki dishes 24 hours prior to the adherence assay to allow formation of a confluent monolayer (Geelen et al., 1993, Infect. Immun. 61:1538-1543). The bacteria are labelled with fluorescein (Geelen et al., supra), adjusted to a concentration of 5 x 10<sup>7</sup> cfu/ml, and added in a volume of 5 μl to at least 6 wells. After incubation at 37 °C for 30 min, the plates are washed and fixed with PBS/glutaraldehyde 2.5%. Attached bacteria are enumerated visually using a fluorescence microscope, such as Nikon Diaphot Inverted Microscope equipped with epifluorescence.

Since two mechanisms, the cell wall and adhesin proteins, determine adherence of a Gram positive bacterium, in particular S. pneumoniae, to a target cell, it may be important to distinguish whether the mutation to the exported protein that inhibits adherence is a mutation to a protein involved in cell wall synthesis or an adhesin. Mutation of the former would have an indirect affect on adherence, while mutation of the latter would directly affect adherence. The following assays can be used to distinguish whether the mutated protein is an adhesin or not: (1) since adherence to macrophages is mainly mediated by exported proteins, adherence assays on macrophages will immediately indicate whether the mutation is to an adhesin; (2) there will be a minimal effect on adherence if bacterial cell wall is separately added in the adherence assay if the mutation is to a protein indirectly involved in adherence, and a further inhibition of adherence if added to a mutant mutated at an adhesin; (3) pretreatment of the bacteria with a protease, such as trypsin, will result in further inhibition of adherence if the mutation is to a protein indirectly involved in adherence, but will have no effect if the mutated protein is an adhesin; and thesin;

(4) once the full length exp gene is isolated, the putative adhesin can be expressed in E. coli or another cell type, or the purified putative adhesin can be covalently associated with different support such as a bacteria, an erythrocyte or an agarose bead, and the ability of the putative adhesin to mediated adherence can be evaluated; (5) the cell wall structure of mutants can be evaluated using standard techniques, in particular HPLC fingerprinting, to determine if the mutation resulted in changes to the cell wall structure, which is indicative of a mutation to a protein indirectly involved with adherence.

10 In another embodiment, the invention provides for identifying genes encoding exported virulence determinants. Generally, virulence determinants can be identified by testing the mutant strain in an animal model for virulence, for example by evaluation of the LD<sub>50</sub> of the animal infected with the strain. An increase in the LD<sub>50</sub> is indicative of a loss of virulence, and therefore the mutation occurred in a locus required for virulence.

The invention also provides for identification of an Exp that is an antigen common to all or many strains of a species of bacterium, or to closely related species of bacteria. This is readily accomplished using an antibody specific to an Exp (the 20 preparation of which is described in detail infra). The ability of the antibody to that particular strain and to all or many other strains of that species, or to closely related species, demonstrates that the Exp is a common antigen. This antibody assay is particularly preferred since it is more immunologically relevant, since the Exp that is a common antigen is an attractive vaccine candidate.

25

Generally, the invention also provides for identification of a functional property of a protein produced by an *exp* gene by comparing the homology of the deduced amino acid or nucleotide sequence to the amino acid sequence of a known protein, or the nucleotide sequence of the gene encoding the protein.

30

Any Gram positive bacterial cell can potentially serve as the nucleic acid source

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for the molecular cloning of an exp gene. The nucleic acid sequences can be isolated from Streptococcus, Bacillus, Mycobacterium, Staphylococcus, Enterococcus, and other Gram positive bacterial sources, etc. The DNA may be obtained by standard procedures known in the art from cloned DNA (e.g., a DNA 5 "library"), by chemical synthesis, by cDNA cloning, or by the cloning of genomic DNA, or fragments thereof, purified from the desired cell (See, for example, Sambrook et al., 1989, supra; Glover, D.M. (ed.), 1985, DNA Cloning: A Practical Approach, MRL Press, Ltd., Oxford, U.K. Vol. I, II). Whatever the source, the gene should be molecularly cloned into a suitable vector for 0 propagation of the gene.

In the molecular cloning of the gene from genomic DNA, DNA fragments are generated, some of which will encode the desired gene. The DNA may be cleaved at specific sites using various restriction enzymes. Alternatively, one may use DNAse in the presence of manganese to fragment the DNA, or the DNA can be physically sheared, as for example, by sonication. The linear DNA fragments can then be separated according to size by standard techniques, including but not limited to, agarose and polyacrylamide gel electrophoresis and column chromatography.

20

Once the DNA fragments are generated, identification of the specific DNA fragment containing the desired exp gene may be accomplished in a number of ways. For example, if an amount of a portion of an exp gene or a fragment thereof is available and can be purified and labeled, the generated DNA fragments may be screened by nucleic acid hybridization to the labeled probe (Benton and Davis, 1977, Science 196:180; Grunstein and Hogness, 1975, Proc. Natl. Acad. Sci. U.S.A. 72:3961). Those DNA fragments with substantial homology to the probe will hybridize. The present invention provides specific examples of DNA fragments that can be used as hybridization probes for pneumococcal exported proteins. These DNA probes can be based, for example, on SEQ ID NOS. 1, 3, 5, 7, 9, 11, 13, 15, 17, 19 or 21. Alternatively, the screening technique of the

invention can be used to isolate additional exp gene fragments for use as probes.

It is also possible to identify the appropriate fragment by restriction enzyme digestion(s) and comparison of fragment sizes with those expected according to a 5 known restriction map if such is available. Further selection can be carried out on the basis of the properties of the gene.

As described above, the presence of the gene may be detected by assays based on the physical, chemical, or immunological properties of its expressed product. For 10 example DNA clones that produce a protein that, e.g., has similar or identical electrophoretic migration, isoelectric focusing behavior, proteolytic digestion maps, proteolytic activity, antigenic properties, or functional properties, especially adhesion activity, as known (or in the case of an adhesion associated protein, unknown) for a particular Exp. In a specific example, infra, the ability of a pneumococcal Exp protein to mediate adhesion is demonstrated by inhibition of adhesion when the protein is mutated. Expression of Exp in another species, such as E. coli. can directly demonstrate whether the exp encodes an adhesin.

Alternatives to isolating the exp genomic DNA include, but are not limited to,

20 chemically synthesizing the gene sequence itself from a known sequence that
encodes an Exp. For example, DNA cloning of an exp gene can be isolated from
Gram positive bacteria by PCR using degenerate oligonucleotides. Other methods
are possible and within the scope of the invention.

25 The identified and isolated gene can then be inserted into an appropriate cloning vector. A large number of vector-host systems known in the art may be used. Possible vectors include, but are not limited to, plasmids or modified viruses, but the vector system must be compatible with the host cell used. In a preferred aspect of the invention, the exp coding sequence is inserted in an E. coli cloning 30 vector. Other examples of vectors include, but are not limited to, bacteriophages such as lambda derivatives, or plasmids such as pBR322 derivatives or pUC

plasmid derivatives, e.g., pGEX vectors, pmal-c, pFLAG, etc. The insertion into a cloning vector can, for example, be accomplished by ligating the DNA fragment into a cloning vector which has complementary cohesive termini. However, if the complementary restriction sites used to fragment the DNA are not present in the cloning vector, the ends of the DNA molecules may be enzymatically modified. Alternatively, any site desired may be produced by ligating nucleotide sequences (linkers) onto the DNA termini; these ligated linkers may comprise specific chemically synthesized oligonucleotides encoding restriction endonuclease recognition sequences. Recombinant molecules can be introduced into host cells via transformation, transfection, infection, electroporation, etc., so that many copies of the gene sequence are generated.

In an alternative method, the desired gene may be identified and isolated after insertion into a suitable cloning vector in a "shot gun" approach. Enrichment for the desired gene, for example, by size fractionation, can be done before insertion into the cloning vector.

In specific embodiments, transformation of host cells with recombinant DNA molecules that incorporate the isolated exp gene or synthesized DNA sequence
20 enables generation of multiple copies of the gene. Thus, the gene may be obtained in large quantities by growing transformants, isolating the recombinant DNA molecules from the transformants and, when necessary, retrieving the inserted gene from the isolated recombinant DNA.

25 The present invention also relates to vectors containing genes encoding analogs and derivatives of Exp's that have the same functional activity as an Exp. The production and use of derivatives and analogs related to an Exp are within the scope of the present invention. In a specific embodiment, the derivative or analog is functionally active, i.e., capable of exhibiting one or more functional activities associated with a full-length, wild-type Exp. As one example, such derivatives or analogs demonstrate adhesin activity.

In particular, Exp derivatives can be made by altering encoding nucleic acid sequences by substitutions, additions or deletions that provide for functionally equivalent molecules. Due to the degeneracy of nucleotide coding sequences, other DNA sequences which encode substantially the same amino acid sequence as 5 an exp gene may be used in the practice of the present invention. These include but are not limited to nucleotide sequences comprising all or portions of exp genes that are altered by the substitution of different codons that encode the same amino acid residue within the sequence, thus producing a silent change. Likewise, the Exp derivatives of the invention include, but are not limited to, those containing. as a primary amino acid sequence, all or part of the amino acid sequence of an Exp including altered sequences in which functionally equivalent amino acid residues are substituted for residues within the sequence resulting in a conservative amino acid substitution. For example, one or more amino acid residues within the sequence can be substituted by another amino acid of a similar polarity, which acts 15 as a functional equivalent, resulting in a silent alteration. Substitutes for an amino acid within the sequence may be selected from other members of the class to which the amino acid belongs. For example, the nonpolar (hydrophobic) amino acids include alanine, leucine, isoleucine, valine, proline, phenylalanine, tryptophan and methionine. The polar neutral amino acids include glycine, serine, 20 threonine, cysteine, tyrosine, asparagine, and glutamine. The positively charged (basic) amino acids include arginine, lysine and histidine. The negatively charged (acidic) amino acids include aspartic acid and glutamic acid.

The genes encoding Exp derivatives and analogs of the invention can be produced

by various methods known in the art. The manipulations which result in their

production can occur at the gene or protein level. For example, a cloned exp gene
sequence can be modified by any of numerous strategies known in the art

(Sambrook et al., 1989, supra). The sequence can be cleaved at appropriate sites
with restriction endonuclease(s), followed by further enzymatic modification if

desired, isolated, and ligated in vitro. In the production of the gene encoding a

derivative or analog of Exp, care should be taken to ensure that the modified gene

remains within the same translational reading frame as the exp gene, uninterrupted by translational stop signals, in the gene region where the desired activity is encoded.

- 5 Additionally, the exp nucleic acid sequence can be mutated in vitro or in vivo, to create and/or destroy translation, initiation, and/or termination sequences, or to create variations in coding regions and/or form new restriction endonuclease sites or destroy preexisting ones, to facilitate further in vitro modification. Any technique for mutagenesis known in the art can be used, including but not limited to, in vitro site-directed mutagenesis (Hutchinson, C., et al., 1978, J. Biol. Chem. 253:6551; Zoller and Smith, 1984, DNA 3:479-488; Oliphant et al., 1986, Gene 44:177; Hutchinson et al., 1986, Proc. Natl. Acad. Sci. U.S.A. 83:710), use of TAB® linkers (Pharmacia), etc. PCR techniques are preferred for site directed mutagenesis (see Higuchi, 1989, "Using PCR to Engineer DNA", in PCR
- 15 Technology: Principles and Applications for DNA Amplification, H. Erlich, ed., Stockton Press, Chapter 6, pp. 61-70).

#### Expression of an Exported Protein

- 20 The gene coding for an Exp, or a functionally active fragment or other derivative thereof, can be inserted into an appropriate expression vector, i.e., a vector which contains the necessary elements for the transcription and translation of the inserted protein-coding sequence. An expression vector also preferably includes a replication origin. The necessary transcriptional and translational signals can also
- 25 be supplied by the native exp gene and/or its flanking regions. A variety of host-vector systems may be utilized to express the protein-coding sequence.
  Preferably, however, a bacterial expression system is used to provide for high level expression of the protein with a higher probability of the native conformation. Potential host-vector systems include but are not limited to
  30 mammalian cell systems infected with virus (e.g., vaccinia virus, adenovirus,
  - etc.); insect cell systems infected with virus (e.g., baculovirus); microorganisms

such as yeast containing yeast vectors, or bacteria transformed with bacteriophage, DNA, plasmid DNA, or cosmid DNA. The expression elements of vectors vary in their strengths and specificities. Depending on the host-vector system utilized, any one of a number of suitable transcription and translation elements may be used.

Preferably, the periplasmic form of the Exp (containing a signal sequence) is produced for export of the protein to the *Escherichia coli* periplasm or in an expression system based on *Bacillus subtillis*. Export to the periplasm can 10 promote proper folding of the expressed protein.

Any of the methods previously described for the insertion of DNA fragments into a vector may be used to construct expression vectors containing a chimeric gene consisting of appropriate transcriptional/translational control signals and the protein coding sequences. These methods may include *in vitro* recombinant DNA and synthetic techniques and *in vivo* recombinants (genetic recombination).

Expression of nucleic acid sequence encoding an exported protein or peptide fragment may be regulated by a second nucleic acid sequence so that the exported 20 protein or peptide is expressed in a host transformed with the recombinant DNA molecule. For example, expression of an exported protein may be controlled by any promoter/enhancer element known in the art, but these regulatory elements must be functional in the host selected for expression. For expression in bacteria, bacterial promoters are required. Eukaryotic viral or eukaryotic promoters, including tissue specific promoters, are preferred when a vector containing an exp gene is injected directly into a subject for transient expression, resulting in heterologous protection against bacterial infection, as described in detail below. Promoters which may be used to control exp gene expression include, but are not limited to, the SV40 early promoter region (Benoist and Chambon, 1981, Nature 30 290:304-310), the promoter contained in the 3' long terminal repeat of Rous sarcoma virus (Yamamoto, et al., 1980, Cell 22:787-797), the herpes thymidine

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kinase promoter (Wagner et al., 1981, Proc. Natl. Acad. Sci. U.S.A. 78:1441-1445), the regulatory sequences of the metallothionein gene (Brinster et al., 1982. Nature 296:39-42); prokaryotic expression vectors such as the  $\beta$ -lactamase promoter (Villa-Kamaroff, et al., 1978, Proc. Natl. Acad. Sci. U.S.A. 75:3727-5 3731), or the tac promoter (DeBoer, et al., 1983, Proc. Natl. Acad. Sci. U.S.A. 80:21-25); see also "Useful proteins from recombinant bacteria" in Scientific American, 1980, 242:74-94; and the following animal transcriptional control regions, which exhibit tissue specificity and have been utilized in transgenic animals: elastase I gene control region which is active in pancreatic acinar cells 10 (Swift et al., 1984, Cell 38:639-646; Ornitz et al., 1986, Cold Spring Harbor Symp. Quant. Biol. 50:399-409; MacDonald, 1987, Hepatology 7:425-515); insulin gene control region which is active in pancreatic beta cells (Hanahan, 1985, Nature 315:115-122), immunoglobulin gene control region which is active in lymphoid cells (Grosschedl et al., 1984, Cell 38:647-658; Adames et al., 1985, 15 Nature 318:533-538; Alexander et al., 1987, Mol. Cell. Biol. 7:1436-1444). mouse mammary tumor virus control region which is active in testicular, breast, lymphoid and mast cells (Leder et al., 1986, Cell 45:485-495), albumin gene control region which is active in liver (Pinkert et al., 1987, Genes and Devel. 1:268-276), alpha-fetoprotein gene control region which is active in liver 20 (Krumlauf et al., 1985, Mol. Cell. Biol. 5:1639-1648; Hammer et al., 1987. Science 235:53-58), alpha 1-antitrypsin gene control region which is active in the liver (Kelsev et al., 1987, Genes and Devel, 1:161-171), beta-globin gene control region which is active in myeloid cells (Mogram et al., 1985, Nature 315:338-340; Kollias et al., 1986, Cell 46:89-94), myelin basic protein gene control region 25 which is active in oligodendrocyte cells in the brain (Readhead et al., 1987, Cell 48:703-712), myosin light chain-2 gene control region which is active in skeletal muscle (Sani, 1985, Nature 314;283-286), and gonadotropic releasing hormone gene control region which is active in the hypothalamus (Mason et al., 1986).

30

Science 234:1372-1378).

Expression vectors containing exp gene inserts can be identified by four general

approaches: (a) PCR amplification of the desired plasmid DNA or specific mRNA. (b) nucleic acid hybridization, (c) presence or absence of "marker" gene functions. and (d) expression of inserted sequences. In the first approach, the nucleic acids can be amplified by PCR with incorporation of radionucleotides or stained with 5 ethidium bromide to provide for detection of the amplified product. In the second approach, the presence of a foreign gene inserted in an expression vector can be detected by nucleic acid hybridization using probes comprising sequences that are homologous to an inserted exp gene. In the third approach, the recombinant vector/host system can be identified and selected based upon the presence or absence of certain "marker" gene functions (e.g., β-galactosidase activity, PhoA activity, thymidine kinase activity, resistance to antibiotics, transformation phenotype, occlusion body formation in baculovirus, etc.) caused by the insertion of foreign genes in the vector. If the exp gene is inserted within the marker gene sequence of the vector, recombinants containing the exp insert can be identified by 15 the absence of the marker gene function. In the fourth approach, recombinant expression vectors can be identified by assaying for the activity of the exp gene product expressed by the recombinant. Such assays can be based, for example, on the physical or functional properties of the exp gene product in in vitro assay systems, e.g., adherence to a target cell or binding with an antibody to the 20 exported protein.

Once a suitable host system and growth conditions are established, recombinant expression vectors can be propagated and prepared in quantity. As previously explained, the expression vectors which can be used include, but are not limited to, the following vectors or their derivatives: human or animal viruses such as vaccinia virus or adenovirus; insect viruses such as baculovirus; yeast vectors; bacteriophage vectors (e.g., lambda), and plasmid and cosmid DNA vectors, to name but a few. The choice of vector will depend on the desired use of the vector, e.g., for expression of the protein in prokaryotic or eukaryotic cells, or as a nucleic acid-based vaccine.

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In addition, a host cell strain may be chosen which modulates the expression of the inserted sequences, or modifies and processes the gene product in the specific fashion desired. Expression from certain promoters can be elevated in the presence of certain inducers; thus, expression of the genetically engineered exported protein may be controlled. Furthermore, different host cells have characteristic and specific mechanisms for the translational and post-translational processing and modification (e.g., cleavage of signal sequence) of proteins. Appropriate cell lines or host systems can be chosen to ensure the desired modification and processing of the foreign protein expressed. Different vector/host expression systems may effect processing reactions, such as proteolytic cleavages, to a different extent.

# Preparation of Antibodies to Exported Proteins

15 According to the invention, recombinant Exp, and fragments or other derivatives or analogs thereof, or cells expressing the foregoing may be used as an immunogen to generate antibodies which recognize the Exp. Such antibodies include but are not limited to polyclonal, monoclonal, chimeric, single chain, Fab fragments, and an Fab expression library.

20

Various procedures known in the art may be used for the production of polyclonal antibodies to a recombinant Exp or derivative or analog thereof. For the production of antibody, various host animals can be immunized by injection with the recombinant Exp, or a derivative (e.g., fragment) thereof, including but not limited to rabbits, mice, rats, etc. In one embodiment, the recombinant Exp or fragment thereof can be conjugated to an immunogenic carrier, e.g., bovine serum albumin (BSA) or keyhole limpet hemocyanin (KLH). Various adjuvants may be used to increase the immunological response, depending on the host species, including but not limited to Freund's (complete and incomplete), mineral gels such as aluminum hydroxide, surface active substances such as lysolecithin, pluronic polyols, polyanions, peptides, oil emulsions, keyhole limpet hemocyanins,

dinitrophenol, and potentially useful human adjuvants such as BCG (bacille Calmette-Guerin) and Corynebacterium parvum.

For preparation of monoclonal antibodies directed toward an Exp or analog

5 thereof, any technique which provides for the production of antibody molecules by
continuous cell lines in culture may be used. These include but are not limited to
the hybridoma technique originally developed by Kohler and Milstein (1975,
Nature 256:495-497), as well as the trioma technique, the human B-cell hybridoma
technique (Kozbor et al., 1983, Immunology Today 4:72), and the EBV
10 hybridoma technique to produce human monoclonal antibodies (Cole et al., 1985,
in Monoclonal Antibodies and Cancer Therapy, Alan R. Liss, Inc., pp. 77-96). In
an additional embodiment of the invention, monoclonal antibodies can be produced
in germ-free animals utilizing recent technology (PCT/US90/02545). According to

the invention, human antibodies may be used and can be obtained by using human hybridomas (Cote et al., 1983, Proc. Natl. Acad. Sci. U.S.A. 80:2026-2030) or by transforming human B cells with EBV virus in vitro (Cole et al., 1985, in Monoclonal Antibodies and Cancer Therapy, Alan R. Liss, pp. 77-96). In fact, according to the invention, techniques developed for the production of "chimeric antibodies" (Morrison et al., 1984, J. Bacteriol. 159-870; Newberger et al., 1984.

Nature 312:604-608; Takeda et al., 1985, Nature 314:452-454) by splicing the genes from a mouse antibody molecule specific for an Exp together with genes from a human antibody molecule of appropriate biological activity can be used; such antibodies are within the scope of this invention. Such human or humanized chimeric antibodies are preferred for use in passive immune therapy (described).

25 infra), since the human or humanized antibodies are much less likely than xenogenic antibodies to induce an immune response, in particular an allergic response, themselves.

According to the invention, techniques described for the production of single chain
antibodies (U.S. Patent 4,946,778) can be adapted to produce Exp-specific single
chain antibodies. An additional embodiment of the invention utilizes the

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techniques described for the construction of Fab expression libraries (Huse et al., 1989, Science 246:1275-1281) to allow rapid and easy identification of monoclonal Fab fragments with the desired specificity for an Exp or its derivatives, or analogs.

5

Antibody fragments which contain the idiotype of the antibody molecule can be generated by known techniques. For example, such fragments include but are not limited to: the F(ab')<sub>2</sub> fragment which can be produced by pepsin digestion of the antibody molecule; the Fab' fragments which can be generated by reducing the disulfide bridges of the F(ab')<sub>2</sub> fragment, and the Fab fragments which can be generated by treating the antibody molecule with papain and a reducing agent.

In the production of antibodies, screening for the desired antibody can be

accomplished by techniques known in the art, e.g., radioimmunoassay, ELISA

15 (enzyme-linked immunosorbant assay), "sandwich" immunoassays, immunoradiometric assays, gel diffusion precipitin reactions, immunodiffusion assays, in situ immunoassays (using colloidal gold, enzyme or radioisotope labels, for example), western blots, precipitation reactions, agglutination assays (e.g., gel agglutination assays, hemagglutination assays), complement fixation assays, immunofluorescence assays, protein A assays, and immunoelectrophoresis assays, etc. In one embodiment, antibody binding is detected by detecting a label on the primary antibody. In another embodiment, the primary antibody is detected by detecting binding of a secondary antibody or reagent to the primary antibody. In a

further embodiment, the secondary antibody is labeled. Many means are known in
the art for detecting binding in an immunoassay and are within the scope of the
present invention. For example, to select antibodies which recognize a specific
epitope of an Exp, one may assay generated hybridomas for a product which binds
to a Exp fragment containing such epitope. For selection of an antibody specific
to an Exp from a particular strain of bacterium, one can select on the basis of
positive binding to that particular strain of bacterium and a lack of binding to Exp
another strain. For selecting an antibody specific to an Exp that is an antigen

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common to all or many strains of a particular bacterium, or to closely related species of bacteria, one can select on the basis of binding to that particular strain and to all or many other strains of that species, or to closely related species.

5 The foregoing antibodies can be used in methods known in the art relating to the localization and activity of Exp, e.g., for Western blotting, imaging Exp, measuring levels thereof in appropriate physiological samples, etc.

# Vaccination and Passive Immune Therapy

10

Active immunity against Gram positive bacteria can be induced by immunization (vaccination) with an immunogenic amount of an exported protein, or an antigenic derivative or fragment thereof, and an adjuvant, wherein the exported protein, or antigenic derivative or fragment thereof, is the antigenic component of the vaccine. Preferably, the protein is conjugated to the carbohydrate capsule or capsules of one or more species of Gram positive bacterium. Covalent conjugation of a protein to a carbohydrate is well known in the art. Generally, the conjugation can proceed via a carbodiimide condensation reaction.

20 The exported protein alone or conjugated to a capsule or capsules cannot cause bacterial infection, and the active immunity elicited by vaccination with the protein according to the present invention can result in both an immediate immune response and in immunological memory, and thus provide long-term protection against infection by the bacterium. The exported proteins of the present invention, or antigenic fragments thereof, can be prepared in an admixture with an adjuvant to prepare a vaccine. Preferably, the exported protein, or derivative or fragment thereof, used as the antigenic component of the vaccine is an adhesin. More preferably, the exported protein, or derivative or fragment thereof, used as the antigenic component of the vaccine is an antigen common to all or many strains of a species of Gram positive bacteria, or common to closely related species of bacteria. Most preferably, the antigenic component of the vaccine is an adhesin

that is a common antigen.

Selection of an adjuvant depends on the subject to be vaccinated. Preferably, a pharmaceutically acceptable adjuvant is used. For example, a vaccine for a human should avoid oil or hydrocarbon emulsion adjuvants, including complete and incomplete Freund's adjuvant. One example of an adjuvant suitable for use with humans is alum (alumina gel). A vaccine for an animal, however, may contain adjuvants not appropriate for use with humans.

- 10 An alternative to a traditional vaccine comprising an antigen and an adjuvant involves the direct in vivo introduction of DNA encoding the antigen into tissues of a subject for expression of the antigen by the cells of the subject's tissue. Such vaccines are termed herein "nucleic acid-based vaccines." Since the exp gene by definition contains a signal sequence, expression of the gene in cells of the tissue
- 15 results in secretion of membrane association of the expressed protein. Alternatively, the expression vector can be engineered to contain an autologous signal sequence instead of the exp signal sequence. For example, a naked DNA vector (see, e.g., Ulmer et al., 1993, Science 259:1745-1749), a DNA vector transporter (e.g., Wu et al., 1992, J. Biol. Chem. 267:963-967; Wu and Wu,
- 20 1988, J. Biol. Chem. 263:14621-14624; Hartmut et al., Canadian Patent Application No. 2,012,311, filed March 15, 1990), or a viral vector containing the desired exp gene can be injected into tissue. Suitable viral vectors include retroviruses that are packaged in cells with amphotropic host range (see Miller, 1990. Human Gene Ther. 1:5-14: Ausubel et al., Current Protocols in Molecular.
- 25 Biology, § 9), and attenuated or defective DNA virus, such as but not limited to herpes simplex virus (HSV) (see, e.g., Kaplitt et al., 1991, Molec. Cell. Neurosci. 2:320-330), papillomavirus, Epstein Barr virus (EBV), adenovirus (see, e.g., Stratford-Perricaudet et al., 1992, J. Clin. Invest. 90:626-630), adeno-associated virus (AAV) (see, e.g., Samulski et al., 1987, J. Virol. 61:3096-3101;
- 30 Samulski et al., 1989, J. Virol. 63:3822-3828), and the like. Defective viruses, which entirely or almost entirely lack viral genes, are preferred. Defective virus

- 40 -

is not infective after introduction into a cell.

Vectors containing the nucleic acid-based vaccine of the invention can be introduced into the desired host by methods known in the art, e.g., transfection, selectroporation, microinjection, transduction, cell fusion, DEAE dextran, calcium phosphate precipitation, lipofection (lysosome fusion), use of a gene gun, or a DNA vector transporter (see, e.g., Wu et al., 1992, J. Biol. Chem. 267:963-967; Wu and Wu, 1988, J. Biol. Chem. 263:14621-14624; Hartmut et al., Canadian Patent Application No. 2,012,311, filed March 15, 1990).

10

Either vaccine of the invention, i.e., a vaccines comprising an Exp antigen or antigenic derivative or fragment thereof, or an exp nucleic acid vaccine, can be administered via any parenteral route, including but not limited to intramuscular, intraperitoneal, intravenous, and the like. Preferably, since the desired result of vaccination is to elucidate an immune response to the antigen, and thereby to the pathogenic organism, administration directly, or by targeting or choice of a viral vector, indirectly, to lymphoid tissues, e.g., lymph nodes or spleen. Since immune cells are continually replicating, they are ideal target for retroviral vector-based nucleic acid vaccines, since retroviruses require replicating cells.

10

Passive immunity can be conferred to an animal subject suspected of suffering an infection with a Gram negative bacterium by administering antiserum, polyclonal antibodies, or a neutralizing monoclonal antibody against the Gram positive bacterium to the patient. Although passive immunity does not confer long term protection, it can be a valuable tool for the treatment of a bacterial infection of a subject who has not been vaccinated. Passive immunity is particularly important for the treatment of antibiotic resistant strains of Gram positive bacteria, since no other therapy is available. Preferably, the antibodies administered for passive immune therapy are autologous antibodies. For example, if the subject is a 20 human, preferably the antibodies are of human origin or have been "humanized," in order to minimize the possibility of an immune response against the antibodies.

An analogous therapy to passive immunization is administration of an amount of an exported protein adhesin sufficient to inhibit adhesion of the bacterium to its target cell. The required amount can be determined by one of ordinary skill using standard techniques.

The active or passive vaccines of the invention, or the administration of an adhesin, can be used to protect an animal subject from infection of a Gram

30 positive bacteria. Thus, a vaccine of the invention can be used in birds, such as chickens turkeys, and pets: in mammals, preferably a human, although the

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vaccines of the invention are contemplated for use in other mammalian species, including but not limited to domesticated animals (canine and feline); farm animals (bovine, ovine, equine, caprine, porcine, and the like); rodents; and undomesticated animals.

5

#### Diagnosis of a Gram Positive Bacterial Infection

The antibodies of the present invention that can be generated against the exported proteins from Gram positive bacteria are valuable reagents for the diagnosis of an 10 infection with a Gram positive bacteriams. Presently, diagnosis of infection with a Gram positive bacterium is difficult. According to the invention, the presence of Gram positive bacteria in a sample from a subject suspected of having an infection with a Gram positive bacterium can be detected by detecting binding of an antibody to an exported protein to bacteria in or from the sample. In one 15 aspect of the invention, the antibody can be specific for a unique strain or a limited number of strains of the bacterium, thus allowing for diagnosis of infection with that particular strain (or strains). Alternatively, the antibody can be specific for many or all strains of a bacterium, thus allowing for diagnosis of infection with that species.

20

Diagnosis of infection with a Gram positive bacterium can use any immunoassay format known in the art, as desired. Many possible immunoassay formats are described in the section entitled "Preparation of Antibodies to Exported Proteins." The antibodies can be labeled for detection in vitro, e.g., with labels such as enzymes, fluorophores, chromophores, radioisotopes, dyes, colloidal gold, latex particles, and chemiluminescent agents. Alternatively, the antibodies can be labeled for detection in vivo, e.g., with radioisotopes (preferably technetium or iodine); magnetic resonance shift reagents (such as gadolinium and manganese); or radio-opaque reagents.

30

Alternatively, the nucleic acids and sequences thereof of the invention can be used

in the diagnosis of infection with a Gram positive bacterium. For example, the 
exp genes or hybridizable fragments thereof can be used for in situ hybridization 
with a sample from a subject suspected of harboring an infection of Gram positive 
bacteria. In another embodiment, specific gene segments of a Gram positive 
bacterium can be identified using PCR amplification with probes based on the exp 
genes of the invention. In one aspect of the invention, the hybridization with a 
probe or with the PCR primers can be performed under stringent conditions, or 
with a sequence specific for a unique strain or a limited number of strains of the 
bacterium, or both, thus allowing for diagnosis of infection with that particular 
10 strain (or strains). Alternatively, the hybridization can be under less stringent 
conditions, or the sequence may be homologous in any or all strains of a 
bacterium, thus allowing for diagnosis of infection with that species.

The present invention will be better understood from a review of the following

15 illustrative description presenting the details of the constructs and procedures that
were followed in its development and validation.

# EXAMPLE 1: GENETIC IDENTIFICATION OF EXPORTED PROTEINS IN STREPTOCOCCUS PNEUMONIAE

20

A strategy was developed to mutate and genetically identify exported proteins in 
Streptococcus pneumoniae. Coupling the technique of mutagenesis with gene 
fusions to phoA, we have developed a tool for the mutation and genetic 
identification of exported proteins from S. pneumoniae. Vectors were created and 
used to screen pneumococcal DNA in Escherichia coli and S. pneumoniae for 
translational gene fusions to alkaline phosphatase (PhoA). In this study the 
identification of several genetic loci that encode exported proteins is reported. By 
similarity to the derived sequences from other genes from prokaryotic organisms 
these loci probably encode proteins that play a role in signal transduction,

macromolecular transport and assembly, maintaining an intracellular chemiosmotic 
balance and nutrient acquisition.

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Twenty five PhoA\* pneumococcal mutants were isolated and the loci from eight of these mutants showed similarity to known exported or membrane associated proteins. Homologs were found to: 1] protein dependent peptide permeases, 2] penicillin binding proteins, 3] Clp proteases, 4] two component sensor regulators,

- 5 5] the phosphoenolpyruvate:carbohydrate phosphotransferase permeases, 6] membrane associated dehydrogenases, 7] P-type (E<sub>1</sub>E<sub>2</sub>-type) cation transport ATPases, 8] ABC transporters responsible for the translocation of the RTX class of bacterial toxins. Unexpectedly one PhoA\* mutant contained a fusion to a member of the D-E-A-D protein family of ATP-dependent RNA helicases
- 10 suggesting export of these proteins.

# Materials and Methods

#### Strains and media.

- 15 The parent strain of S. pneumoniae used in these studies was R6x, which is a derivative of the unencapsulated Rockefeller University strain R36A (Tiraby and Fox, 1973, Proc. Natl. Acad. Sci. U.S.A. 70:3541-3545). E. coli strains used were DH5α, which is F f80dlacZ Δ(lacZYAΔM15) lacU169 recA1 endA1 hsdR17 (fx-mx,) supE44 I thy-1 gyrA relA1 (Bethesda Research Laboratories): CC118.
- 20 which is Δ(ara leu)7697 ΔlacX74 araD139 phoA20 galE galK thi rpsE rpoB argE recA1 (Manoil and Beckwith, 1985, Proc. Natl. Acad. Sci. U.S.A. 82:8129-8133), S1179 which is F ΔlacU169 dam3 rpsL (Brown, 1987, Cell. 49:825-33); and JCB607, which contains an expression vector for the production DsbA (rna met pBJ41 pMS421) (Bardwell et al., 1991, Cell. 67:581-589). Strains of S.
- 25 pneumoniae and their relevant characteristics generated in this study are listed in Table 1.

- 45 -

Table 1. Bacterial strains of Streptococcus pneumoniae created in this study

	Strain	Relevant characteristics	Gene Family or Homolog	Source
	R6x	Hex, Parent strain		(Tiraby and Fox, 1973)
5	SPRU2	PhoA fusion to signal sequence 1		Current study
	SPRU37	PhoA fusion to signal sequence 2		Current study
	SPRU96	pHRM100::222		Current study
	SPRU97	pHRM104::222		Corrent study
	SPRU121	PhoA fusion to AmiA	peptide permeases	Current study
0	SPRU98	PhoA fusion to Expl	peptide permeases	Current study
	SPRU42	PhoA fusion to Exp2 (PonA)	penicillin binding protein 1a	Current study
	SPRU40	PhoA fusion to Exp3	two component family of sensor regulators	Current study
	SPRU39	PhoA fusion to Exp4	Clp proteases	Current study
	SPRU87	PhoA fusion to Exp5	PTS family of permeases	Current study
5	SPRU24	PhoA fusion to Exp6	glycerol-3-phosphate dehydrogenase; GlpD; B. subtilis	Current study
	SPRU75	PhoA fusion to Exp7	P-type cation transport ATPases	Current study
	SPRU81	PhoA fusion to Exp8	RTX type traffic ATPases	Current study
	SPRU17	PhoA fusion to Exp9	ATP dependent RNA helicases	Current study

20 The derived amino soid sequences were determined from plasmids recovered from the PhoA\* mutants. Homologs were identified by searching a protein database with the BLAST algorithm. See Figure 5 for all properties.

S. pneumoniae were routinely plated on tryptic soy agar supplemented with sheep blood (TSAB) to a final concentration of 3% (vol./vol.). Cultures were also grown in a liquid semi synthetic casein hydrolysate medium supplemented with yeast extract (C+Y medium) (Lacks and Hotchkiss, 1960, Biochem. Biophys. Acta. 39:508-517). In some instances, S. pneumoniae were grown in Todd Hewitt broth (THBY) supplemented with yeast to a final concentration of 5% (w/v).

30 Where indicated, S. pneumoniae was grown in C+Y in the presence of the

disulfide oxidant 2-hydroxyethyl disulfide at a concentration of 600 µM, which is 5 times less than the minimal inhibitory concentration required for growth. E. coli were grown in either liquid or on solid Luria-Bertani (LB) media. Selection of E. coli with plasmid vectors was achieved with erythromycin (erm) at a concentration of 500 µg / ml. For the selection and maintenance of S. pneumoniae containing chromosomally integrated plasmids, bacteria were grown in the presence of 0.5 to 1 µg / ml of erm.

Transformation of S. pneumoniae was carried out as follows: Bacteria were
10 grown in C+Y medium at 37°C and samples were removed at 10 min. intervals
between an O.D. 600 of 0.07 and 0.15 and stored at -70°C in 10% glycerol.

Samples were thawed on ice and DNA (final concentration, 1 µg / ml) was added
before incubation at 37°C for 90 min. Transformants were identified by selection
on TSAB containing the appropriate antibiotic.

# Recombinant DNA techniques.

15

Plasmids pHRM100 and pHRM104 (Figure 1) were constructed by insertion of either the 2.6 kB Smal or BamHI fragments of pPHO7, which contain the truncated gene for phoA (Guitierrez and Devedjian, 1989, Nucleic Acid Res.

- 20 17:3999), into the corresponding sites in pJCD9 (Chen and Morrison, 1988, Gene. 64:155-164). A unique Smal cloning site for pHRM100 and a unique BamHI cloning site for pHRM104 upstream from phoA were generated by selective deletion of duplicated sites.
- 25 Chromosomal DNA from S. pneumoniae was prepared by the following procedure: Cells were grown in 10 ml of THBY or C+Y with 0.5 μg / ml erm to an O.D.620 of 0.7. The cells were isolated by centrifugation and washed once in 500 μl of TES (0.1 M Tris-HCl, pH 7.5; 0.15 M NaCl, 0.1 M ethylenediaminetetra-acetic acid (EDTA)). The supernatant was discarded and the 30 pellet resuspended in 500 μl of fresh TES. Bacteria were lysed with the addition of 50 μl of 1% (vol./vol.) deoxycholate. The lysate was sequentially incubated

with RNase (2 μg) and pronase (400 ng) for 10 min. at 37 °C. This solution was extracted three times with an equal volume of phenol:chloroform:isoamyl alcohol (25:24:1), followed by one extraction with an equal volume of chloroform:isoamyl alcohol (24:1). The DNA was precipitated with the addition of two volumes of cold ethanol, washed once with 70% ethanol, and resuspended in 10 mM Tris-HCl, pH 8.0, 1 mM EDTA. In some instances this protocol was adjusted to accommodate 400 ml of bacteria.

Plasmid libraries containing pneumococcal DNA were created with pHRM100 and pHRM104 in E. coli for insertion duplication mutagenesis in S. pneumoniae.

Chromosomal DNA from S. pneumoniae was digested for 18 hr. with either Alul or Rsal or for 1.5 hr. with Saulila. This DNA was size fractionated on a 0.7% agarose gel and 400-600 base pair fragments were extracted and purified with glass beads (BiO 101 Inc., La Jolla, CA) according to the manufacturer's 15 instructions. DNA was ligated for 18 hr. at 4'C into either the Smal or BamHI sites of pHRM100 or pHRM104, respectively, at insert to vector ratio of 6:1. The ligation mixture was transformed into the E. coli strain S1179 or the PhoA strain CC118. Plasmid DNA was obtained from these libraries using the Qiagen midi plasmid preparation system (Qiagen Inc., Chatsworth, CA) according to the 20 manufacturer's instructions.

The mutagenesis strategy in S. pneumoniae involved insert duplication upon plasmid integration (Figure 1b). Because of this duplication there was a low frequency excision of the integrated plasmid with its insert that contaminated chromosomal preparations of pneumococcal DNA. Therefore, integrated plasmids containing a pneumococcal insert were easily recovered from S. pneumoniae by transformation of these excised plasmids directly into competent E. coli.

To create a gene fusion between the phoA and amiA, a 600 base pair fragment of 0 amiA was obtained by the polymerase chain reaction of chromosomal DNA from S. pneumoniae using the forward and reverse primers:

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5'AAAGGATCCATGAARAARAAYMGHGTNTTY3' (SEO ID NO:40). and

5'TTTGGATCCGTTGGTTTAGCAAAATCGCTT3' (SEQ ID NO:41) respectively, where R=A/G, Y=T/C, M=C/A, H=T/C/A and N=G/A/T/C. 5 Amplification of DNA was carried out with 50 ng of chromosomal DNA, 2 mM of the forward primer, 1 mM of the reverse primer and 2.5 U of AmpliTag DNA polymerase (Perkin Elmer, Norwalk, CT), dNTPs and buffer provided by the manufacturer. Amplification (30 rounds) was carried out using the following procedure: 1 min. at 94 °C for denaturation, 2 min. at 72 °C for extension, and 1 10 min, at 45°C for reannealing. A 600 base pair fragment was obtained, digested with BamHI and ligated into the corresponding site of pHRM104. This mixture was transformed into E. coli and a single recombinant clone that contained the vector with the insert was identified. An inframe coding sequence across the fusion joint was confirmed by sequence analysis. Plasmid DNA from this clone 15 was transformed into S. pneumoniae and transformants were screened for PhoA activity by the colony lift assay to confirm production and export of the fusion protein.

# DNA sequencing.

- 20 Oligonucleotides (5'AATATCGCCCTGAGC3', SEQ ID NO:42; and 5'ATCACGCAGAGCGGCAG3', SEQ ID NO:43) were designed for sequencing across the fusion joints of the pneumococcal inserts into pHRM100 and pHRM104. Double stranded sequence analysis was performed on plasmid DNA by the dideoxy-chain termination method (Sanger et al., 1977, Proc. Natl. Acad.
- 25 Sci. U.S.A. 74:5463-5467) using the Sequenase Version 2.0 DNA sequencing kit (United States Biochemical Corp., Cleveland, Ohio) according to the manufacturer's instructions. Dimethylsulfoxide (1% vol. / vol.) was added to the annealing and extension steps.

#### 30 Alkaline phosphatase activity.

Even though alkaline phosphatase has been characterized in some Gram positive

organisms such as Enterococcus faecalis (Rothschild et al., 1991, In "Genetics and Molecular Biology of Streptococci, Lactococci, and Enterococci.", Dunny, et al., Washington D.C. American Society for Microbiology, pp. 45-48) and B. subtilis (Chesnut et al., 1991, Mol. Microbiol. 5:2181-90; Hulett et al., 1991, J. Biol. Chem. 266:1077-84; Sugahara et al., 1991, J. Bacteriol. 173-1824-6), nothing is known about this enzyme in S. pneumoniae. PhoA activity associated with the parental strain of S. pneumoniae was measured with chromogenic substrates in the assays described below and gave nominal results. Therefore, detection of PhoA activity due to the expression of fusion proteins in S. pneumoniae was performed in a low or negative background.

To screen for pneumococcal derived PhoA fusions in *E. coli*, plasmid libraries were screened in the PhoA strain CC118. Transformants were plated on LB media supplemented with 40 to 80 µg / ml of the chromogenic substrate 5-bromo-15 4-chloro-3-indolyl phosphate (XP). Blue colonies developed in 15 to 24 hr, and indicated PhoA activity. Individual colonies were streak purified on fresh LB/XP plates to verify the blue phenotype.

To screen for PhoA<sup>+</sup> mutants of *S. pneumoniae*, individual colonies were screened in a colony lift assay with XP as adapted from a previously described procedure (Knapp and Mekalanos, 1988, J. Bacteriol. 170:5059-5066). Individual two day old colonies were transferred to nitrocellulose filters (HAHY, Millipore, Bedford, MA) and air dried for two to five min. The filters were placed colony side up on No. 3 filter papers (Whatman, Inc. Clifton, NJ), pre-soaked in 0.14 M NaCl, and incubated for 10 min. at 37°C. This was repeated once and then the membranes were transferred to fresh filter papers pre-soaked in 1 M Tris-HCl, pH 8.0 and incubated for 10 min. at 37°C. Finally the membranes were transferred to another fresh filter paper soaked in 1 M Tris-HCl, pH 8.0, with 200 μg / ml of XP and incubated at 37°C. Blue colonies indicated PhoA<sup>+</sup> mutants and were detected in 10 min. to 18 hr. Colonies were picked either directly from the filters or from the original plates. After colonies were streak purified on TSAB plates, the blue

phenotype was reconfirmed in a subsequent colony lift assay.

PhoA activity expressed in strains of S. pneumoniae was determined from exponentially growing cultures. Bacteria from 10 ml cultures were isolated by
5 centrifugation, washed once in saline and resuspended in 1 ml of 1 M Tris-HCl, pH 8.0. Activity was determined by hydrolysis of p-nitrophenol phosphate in a previously described assay (Brickman and Beckwith, 1975, Mol. Biol. 96:307-316; Guitierrez et al., 1987, J. Mol. Biol. 195:289-297). Total protein was determined on lysed bacteria with Coomassie blue dye (Bradford, 1976, Anal. Biochem.
0 72:248-254).

# Purification of DsbA.

DsbA was purified to near homogeneity from an *E. coli* strain (ICB607) that contains an expression vector with the corresponding gene (Bardwell et al., 1991, 15 Cell. 67:581-589). Briefly, 2 ml of a fresh overnight culture was added to 400 ml of LB media and grown for 2 hr. at 37°C. The culture was adjusted to 3 mM isopropyl β-D-thiogalactopyranoside (IPTG) and grown for an additional 2 hr. Bacteria were isolated by centrifugation and resuspended in 6 ml of 100 mM Tris-HCl pH 7.6, 5 mM EDTA and 0.5 M sucrose. This suspension was incubated for 10 min. on ice and the cells isolated by centrifugation. Bacteria were resuspended in 6 mL of 5 mM MgCl<sub>2</sub> and incubated for 10 min. on ice. The supernatant was isolated after centrifugation. This material contained a predominant Coomassie blue stained band with an apparent M, of 21 kDa on an SDS polyacrylamide gel, which is identical to that of DsbA, and was judged to be approximately 95% pure

#### Subcellular fractionation.

Pneumococci were separated into subcellular fractions by a modification of a previously described technique (Hakenbeck et al., 1986, Antimicrobial agents and chemotherapy. 30:553-558). Briefly, bacteria were grown in 10 ml of C+Y medium to an O. D. 620 of 0.6, and isolated by centrifugation at 17,000xg for 10

min. Cell pellets were resuspended in 250 μl of TEP (25 mM Tris-HCl pH 8.0, 1 mM EDTA, 1 mM phenyl methyl sulfonyl fluoride). The suspension was sonicated for a total of 4 min. with 15 sec. bursts. Greater than 99% of the bacteria were broken as revealed by visual inspection. Cellular debris was
5 removed by centrifugation (17,000xg for 10 min.). The bacterial membranes and the cytoplasmic contents were separated by centrifugation at 98,000 x g for 4 hr in a Beckman airfuge. The supernatant from this final step contained the cytoplasmic fraction while the pellet contained the bacterial membranes. Samples from each fraction were evaluated for protein content and solubilized in SDS sample buffer
10 for subsequent gel electrophoresis.

# Immunological detection of fusion proteins.

Total bacterial lysates and subcellular fractions were subjected to SDSpolyacrylamide gel electrophoresis and proteins transferred to nitrocellulose

15 membranes (Immobilon, Millipore, Bedford, MA) using the PhastSystem
(Pharmacia LKB, Uppsula Sweden) according to the manufacturer's instructions.

The membranes were probed with polyclonal anti-PhoA antibodies (5 Prime - 3
Prime, Boulder, CO) at a dilution of 1:1000, with a peroxidase conjugated second
antibody at a dilution of 1:1000. Immunoreactive bands were detected with
20 hydrogen peroxide and diaminobenzidine or by enhanced chemiluminescence with
chemicals purchased from Amersham (Arlington Heights, IL).

# Results and Discussion

25 Construction of reporter plasmids and pneumococcal libraries.
In order to genetically screen for exported proteins in S. pneumoniae by insertion duplication mutagenesis, a truncated form of phoA (Guitierrez and Devedjian, 1989, Nucleic Acid Res. 17:3999) was placed in the pneumococcal shuttle vector pJDC9 (Figure 1a) (Chen and Morrison, 1988, Gene. 64:155:164) Two vectors were created with either a unique Smal (pHRM100) or a unique BamHI (pHRM104) cloning site 5' to phoA. The cloning sites in each vector are flanked

by two KpnI sites to facilitate easy identification of an insert.

Efficient insertion duplication mutagenesis requires the cloning of an internal gene fragment within the vector prior to integration (Figure 1b). Therefore plasmid 5 libraries were created in *E. coli* with 400 to 600 base pair inserts of pneumococcal DNA. Several libraries representing approximately 2,600 individual clones were screened for translational fusions to phoA in either *E. coli* or *S. pneumoniae*.

# Identification of pneumococcal PhoA fusions in E. coli.

10 When the pneumococcal libraries representing 1,100 independent clones were screened in the PhoA E. coli strain CC118 fifty five colonies displayed the blue phenotype when plated on media containing 5-bromo-4-chloro-3-indolyl phosphate (XP). Since the cloning vectors pHRM100 and pHRM104 do not contain an intrinsic promoter upstream from phoA, fusion proteins derived from these 15 plasmids must have been generated from pneumococcal DNA that contains a promoter, a translational start site and functional signal sequence. DNA sequence analysis of the inserts from two of these plasmids showed a putative promoter, ribosome binding sites and coding sequences for 48 and 52 amino acids that were inframe with the coding sequence for phoA. These coding sequences have features characteristic of prokaryotic signal sequences such as a basic N-terminal region, a central hydrophobic core and a polar C-terminal region (von Heijne, 1990, J. Memb, Biol. 115:195-201) (Table 2).

Table 2. Predicted coding regions from two genetic loci that produced PhoA

25 fusion proteins in both S. pneumoniae and E. coli.

30

Strain	Signal sequence *
SPRU2	MKHLLSYFKPYIKESILAPLFKLLEAVFELLVPMVIA,GIVDQSLPQ GDPRVP (SEQ ID NO:44)
SPRU37	MAKNNKVAVVTTVPSVAEGLKNVNG, VNFDYKDEASAKEAIKEE KLKGYLTIDPRVP (SEO ID NO:45)

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The coding regions were identified from the DNA sequences 5' to phoA from the plasmids recovered from these strains. The arrow indicates the predicted signal peptide cleavage site based on the "-3, -1 rule" (von Heijne, 1986, Nucleic Acid Res. 14:4683-4690) and the amino acids in bold face type are from the coding region for phoA.

A putative cleavage site was identified in both sequences with an algorithm designed to identify such sites based on the "-3, -1 rule" (von Heijne, 1986, Nucleic Acid Res. 14:4683-4690). Transformation and integration of these plasmids into S. pneumoniae gave transformants that produced blue colonies in the colony lift assay and each produced anti-PhoA immunoreactive fusion proteins with an apparent M, of 55 kDa on SDS polyacrylamide gels (data not shown). These results clearly show that heterologous signal sequences from S. pneumoniae fused to PhoA are functional in both E. coli and S. pneumoniae and probably use a similar secretion pathway.

# PhoA fusions to an exported pneumococcal protein.

are responsible for the transport of small peptides (Alloing et al., 1989, Gene. 20 76:363-8; Alloing et al., 1990, Mol. Microbiol. 4:633-44; Gilson et al., 1988, EMBO J. 7:3971-3974). AmiA contains a signal sequence and should be an exported lipoprotein attached to the bacterial membrane by a lipid mojety

AmiA is a pneumococcal representative of the family of bacterial permeases that

exported lipoprotein attached to the bacterial membrane by a lipid moiety covalently linked to the N-terminal cysteine (Gilson et al., 1988, EMBO J. 7:3971-3974). We genetically engineered a pneumococcal mutant (SPRU121) that contained the 5' coding region of amiA fused inframe at codon 169 to phoA.

Colonies of this mutant produced the blue phenotype when exposed to XP suggesting that the hybrid protein was exported. An immunoreactive polypeptide with the predicted M, of 67 kDa was confirmed by Western analysis of a total cell lysate (data not shown).

30

5

# Identification of PhoA fusions in S. pneumoniae.

Encouraged by the detection of PhoA fusions derived from pneumococcal DNA in both E. coli and S. pneumoniae, we created a library of pneumococcal

transformants that contained random chromosomal insertions of the PhoA vectors pHRM100 and pHRM104. From a bank of 1,500 clones, 75 mutants were isolated that displayed the blue phenotype in the colony lift assay with XP. Because S. pneumoniae spontaneously lyse during stationary growth due to an endogenous amidase (LytA), we were concerned that the blue phenotype of some of the mutants was the result of cell lysis and not due to the export of a fusion protein from viable cells. The DNA from 10 random blue mutants that included SPRU22, 42, 75, 81, and 98 was transformed into a lytA minus background and all still displayed the blue phenotype (data not shown).

10

One of the mutants (SPRU98) displayed the blue phenotype on XP and expressed a 93 kDa anti-PhoA immunoreactive polypeptide (Fig 2: lane 2). Since the coding region to phoA would produce a polypeptide with a molecular mass of 49 kDa, we can conclude that the fusion protein was being produced from a coding region corresponding to a polypeptide with a molecular mass of 44 kDa. In contrast, mutants SPRU96 and 97, that contained randomly inserted vectors and were not blue when exposed to XP, did not produce any immunoreactive material (Fig 2: lanes 3, 4). The fusion protein from SPRU98 was proteolytically degraded when whole bacteria were exposed to low concentrations of trypsin suggesting an extracellular location (Fig 2, lane 5). Consistent with this result was the direct measurement of alkaline phosphatase activity associated with whole bacteria. Compared to the parental strain and a PhoA mutant (SPRU97) with a randomly integrated plasmid, there was a three- to four-fold greater enzyme activity for SPRU98 (Table 3). Collectively these results suggest that PhoA fusions to exported proteins were translocated across the cytoplasmic membrane of S. pneumoniae.

Table 3. Alkaline phosphatase activity for a pneumococcal mutant with a gene fusion to phoA.

30

Strain Integrated phoA Colony lift assay b Phosphatase activity c

15

-	SPRU98 +		blue 44.7 ±6		·	
	SPRU97	+	white	18.4 ±5		
5	R6x	0	white	14.6 ±4		

\* SPRU97 and SPRU98 contain the phoA vector pHRM104 randomly integrated into the chromosome as described in the text.

10 The PhoA\* mutant was isolated based on the expression of alkaline phosphatase activity detected by exposure of individual colonies to XP in the colony lift assay.
\*Units of alkaline phosphatase activity were determined as described in Experimental procedures. The assay was performed on washed cells from exponentially growing cultures. The results are presented as units of enzyme activity / mg of total protein.

Disulfide oxidants increase the enzyme activity of PhoA fusions in S. pneumoniae. In E. coli. PhoA activity requires protein translocation across the cytoplasmic membrane, incorporation of Zn2+, disulfide bond formation and dimerization. Following this activation process the enzyme is highly protease resistant (Roberts and 20 Chlebowski, 1984). Recently two groups have identified a single genetic locus, dsh4 (Bardwell et al., 1991, Cell. 67:581-589), and ppfA (Kamitani et al., 1992, EMBO J. 11:57-67), that encodes a disulfide oxidoreductase, which facilitates the formation of disulfide bonds in PhoA. A similar locus has also been identified in V. cholerae (Peek and Taylor, 1992, Proc. Natl. Acad. Sci. 89:6210-6214). Mutations in dsbA dramatically decreased PhoA activity and rendered the protein protease sensitive both in vitro and in vivo (Bardwell et al., 1991, Cell. 67:581-589; Kamitani et al., 1992. EMBO J. 11:57-67). Since the enzyme activity associated with the PhoA fusions in S. pneumoniae was universally 10 fold lower than values obtained with fusions in E. coli (data not shown) and due to the protease sensitivity of the PhoA fusion depicted 30 in Figure 2, we hypothesized that the addition of DsbA or a strong disulfide oxidant would promote disulfide bond formation, increase enzyme activity and retard proteolytic degradation.

SPRU98 which produces a PhoA fusion protein with an M<sub>r</sub> of 93 kDa was grown in 35 either the presence of 10  $\mu$ M DsbA or 600  $\mu$ M 2-hydroxyethel disulfide, a strong disulfide oxidant. Under both conditions enzyme activity was increased at least two fold (Table 4).

5 Table 4. Effect of disulfide oxidants on the alkaline phosphatase activity

	Agent	
10	10 μM DsbA	138.4 ±7
	600 μM 2-hydroxyethel disulfide	107.5 ±8
	Control	51.2 ±5

15

The strain SPRU98 (10 ml) was grown in the presence of the indicated agents to mid log phase  $(OD_{eco}: 0.4)$ , concentrated and assayed for alkaline phosphatase activity. Hydrolysis of p-nitrophenol phosphate was determined with whole bacteria in the presence of 1 M Tris-HCl, pH 8.0 for one hr. at 37  $\blacksquare$ C. Activity units are expressed per mg of total protein.

Compared to the control, there was also an increased amount of immunoreactive protein detected in the presence of these two compounds (Figure 3). This suggested increased protein stability and resistance to intrinsic proteolysis. Since there was only a modest increase in enzyme activity conveyed by these compounds, we propose that there may be other factors required for the correct folding of PhoA that are absent in S. pneumoniae. It is of note that the derived sequences of other alkaline phosphatase isozymes identified in the Gram positive organisms B. subtilis (Chesnut et al., 1991, Mol. Microbiol. 5:2181-90; Hulett et al., 1991, J. Biol. Chem. 30 266:1077-84; Sugahara et al., 1991, J. Bacteriol. 173:1824-6) and Enterococcus faecalis contain only one or no cysteine residues. This may suggest that the presence of an oxido-reductase system for the correct folding of these intra or intermolecular disulfide bonds may be a unique property of some Gram negative organisms which contain a well defined periplasm.

35

Identification of exported proteins by sequence analysis of the PhoA fusions from S. pneumoniae.

The plasmids containing pneumococcal inserts were recovered in E. coli from 48 pneumococcal mutants that displayed the blue phenotype on XP. Digestion of these plasmids with KpnI dissects the pneumococcal inserts from the parent vector. The size of the inserts were all approximately 400 to 900 base pair. Preliminary sequence 5 analysis of the 48 inserts revealed 21 distinct sequences, thus demonstrating a sibling relationship between some of the mutants. Long coding regions corresponding to 50 to 200 amino acids inframe with PhoA were established for most of the inserts, nine of which are presented in Figure 4. Using the BLAST algorithm (Altschul et al., 1990. J. Mol. Biol. 215:403-410), the derived protein sequences were analyzed for 10 similarity to sequences deposited in the most current version of the non redundant protein database at the National Center for Biotechnology Information (Washington. D. C.). Sequence from these nine inserts (Figure 4) revealed coding regions with similarity to families of eight known exported or membrane associated proteins (Figure 5). Those proteins encoded by the genes that correspond to the potential 15 reading frames without a known function are designated with the preface exp (exported protein) to describe the different genetic loci.

No similarity between the derived sequences from the other inserts to those in the data base was detected. The sequences for all nine inserts will be made available in 20 Genbank (Accession numbers: to be assigned) after the filing date of this application.

Exp1 showed similarity to the family of permeases responsible for the transport of small peptides in both Gram negative and Gram positive bacteria (Figure 5A). The reading frame identified showed the greatest similarity to the exported protein, AmiA, from S. pneumoniae (Alloing et al., 1990, Mol. Microbiol. 4:633-44). The ami locus was first characterized in a spontaneous mutant resistant to aminopterin (Sicard, 1964, Genetics. 50:31-44; Sicard and Ephrussi-Taylor, 1965). The wild type allele may be responsible for the intracellular transport of small branched chain amino acids (Sicard, 1964). Exp1 is clearly distinct from AmiA and represents a related member of the family of permeases present in the same bacteria. E. coli has at least three peptide permeases while B. subtilis has at least two (for a review see (Higgins et al.,

1990, J. Bioengen. Biomembranes. 22:571-92)). Mutations in an analogous locus SpoOK from B. subtilis inhibit sporulation and dramatically decrease transformation efficiency in naturally competent cells (Perego et al., 1991, Mol. Microbiol. 5:173-85; Rudner et al., 1991, J. Bacteriol). Recent results have shown that mutations in expl also decrease transformation efficiency in S. pneumoniae whereas mutations in amiA did not. Therefore, two distinct peptide permeases from two different Gram positive bacteria affect the process of transformation in these naturally competent bacteria.

Both the DNA and derived protein sequences of exp2 were identical to ponA (basepairs 1821-2055) which encodes penicillin-binding protein 1A (PBP1a) (Martin et al., 1992a, J. Bacteriol. 174:4517-23) (Figure 5B). This protein belongs to the family of penicillin-interacting serine D, D-peptidases that catalyze the late steps in murein biosynthesis. PBP1a is routinely isolated from pneumococcal membrane
preparations and is generally considered an exported protein (Hakenbeck et al., 1991, J. Infect. Dis. 164:313-9; Hakenbeck et al., 1986, Antimicorbial Agents and Chemotherapy. 30:553-558; Martin et al., 1992, Embo J. 11:3831-6). In E. coli deletions of both PBP1a and PBP1b are lethal to the cell but the bacteria are able to compensate if either gene is deleted (Yousif et al., 1985, J. Gen. Microbiol.
131:2839-2845). It would be interesting to compare the peptidoglycan profile of SPRU42 to the parent strain to determine if the gene fusion to PBP1a alters enzyme function.

Exp3 showed significant sequence similarity to PilB from N. gonorrhoeae (Figure 5C)

(Taha et al., 1988, EMBO J. 7:4367-4378). There were two regions of similarity which correspond to the C-terminal domain of PilB. There was a short gap of 25 amino acids for Exp3 and 37 amino acids for PilB which showed no similarity. This suggests a modular structure function relationship for these two proteins. Consistent with this result, PhoA-PilB hybrids were localized to the membrane fraction of N. gonorrhoeae (Taha et al., 1991, Mol. Microbiol 5:137-48) indicating membrane translocation.

It has been suggested that PilA and PilB are members of the family of two component sensor regulators that control pilin gene expression and that PilB is a transmembrane sensor with the conserved transmitter region that contains kinase activity in the C-terminal region of the protein (Taha et al., 1991, Mol. Microbiol. 5:137-48; Taha et al., 1992, J. Bacteriol. 174:5978-81). The conserved histidine residue (H<sub>408</sub>) in PilB required for autophosphorylation that is characteristic of this family is not present in Exp3. Since no pilin has been identified on S. pneumoniae one would assume a different target site for gene regulation by Exp3.

- 10 The coding region identified with Exp4 suggests that it is similar to the ubiquitous family of Clp proteins found in both eukaryotes and prokaryotes (Figure 5D) (for a review see Squires and Squires, 1992, J. Bacteriol. 174:1081-1085). Exp4 is most similar to the homolog CD4B from tomato (Gottesman et al., 1990, Proc. Natl. Acad. Sci. U.S.A. 87:3513-7) but significant similarity was also noted to ClpA and ClpB from E. coli. It has been proposed that these proteins function either as regulators of proteolysis (Gottesman et al., 1990, Proc. Natl. Acad. Sci. U.S.A. 87:3513-7) or as molecular chaperones (Squires and Squires, 1992, J. Bacteriol. 174:1081-1085). One universal feature of the Clp proteins is a long leader sequence that implies membrane translocation (Squires and Squires, 1992, supra, J. Bacteriol. 174:1081-1085). Indeed, plant ClpC is translocated into chloroplasts (Moare, 1989, Ph.D. thesis. University of Wisconsin, Madison). Even though little is known about the subcellular localization of the other Clp proteins, our results suggest translocation of the pneumococcal homolog across the bacterial membrane.
- 25 Exp5 showed similarity to PtsG from B. subtilis (Gonzy-Tréboul et al., 1991, Mol. Microbiol. 5:1241-1249) which is a member of the family of phosphoenolpyruvate:carbohydrate phosphotransferase permeases that are found in both Gram positive and Gram negative bacteria (for a review see Saier and Reizer, 1992, J. Bacteriol. 174:1433-1448) (Figure 5E). These permeases are polytopic 30 membrane proteins with several translocated domains.

Analysis of the insert recovered from Exp6 revealed a coding region with similarity to glycerol-3-phosphate dehydrogenases from several prokaryotic species (Figure 5F). It is most similar to GlpD from B. subtilis (Holmberg et al., 1990, J. Gen. Microbiol. 136:2367-2375). This enzyme is a membrane associated flavoprotein forming a 5 complex with cytochrome oxidases which are integral membrane proteins. Besides converting glycerol-3-phosphate to dihydroxyacetone phosphate and glyceraldehyde-3phosphate for subsequent entry into the glycolytic pathway, this enzyme delivers electrons to the cytochrome oxidases for subsequent transport. It has been proposed that these dehydrogenases are bound to the inner surface of the cytoplasmic 10 membrane via nonspecific hydrophobic interactions (Halder et al., 1982, Biochemistry, 21:4590-4606; Koland et al., 1984, Biochemistry, 23:445-453; Wood et al., 1984, Biochem. J. 222:519-534). Alternatively it has been proposed that there are a specific and saturable number of binding sites between the dehydrogenases and the cytochromes serving to anchor the dehydrogenases to the cytoplasmic membrane. 15 The data reported here suggest that in S. pneumoniae a segment of the dehydrogenase is translocated to the outer surface of the bacteria (Kung and Henning, 1972, Proc. Natl. Acad. Sci. U.S.A. 69:925-929). Translocation of the catalytic domain would certainly not alter enzyme function. In reconstituted inside out membrane vesicles. electron transfer to the cytochromes occurred when dehydrogenases were added to either side of the vesicles (Halder et al., 1982, Biochemistry, 21:4590-4606).

Analysis of the derived sequence for Exp7 showed similarity to the family of both eukaryotic and prokaryotic P-type (E,E<sub>2</sub>-type) cation transport ATPases responsible for the transport of cations such as Ca<sup>2+</sup>, Mg<sup>2+</sup>, K<sup>+</sup>, Na<sup>+</sup>, and H<sup>+</sup> (Figure 5G).

These ATPases are intrinsic membrane proteins with several translocated domains. Examples have been identified in E. faecalis (Solioz et al., 1987, J. Biol. Chem. 262:7358-7362), Salmonella typhimurium (Snavely et al., 1991, J. Biol. Chem. 266:815-823), E. coli (Hesse et al., 1984, Proc. Natl. Acad. Sci. U.S.A. 81:4746-4750), Neurospora crassa (Addison, 1986, J. Biol. Chem. 26:14896-14901; Hager et al., 1986, Proc. Natl. Acad. Sci. U.S.A. 83:7693-7697), Saccharomyces cerevisiae (Rudolph et al., 1989, Cell. 58:133-145) and the sarcopolasmic reticulum

of rabbit skeletal muscle (Brandi et al., 1986, Cell. 44:597-607; Serrano et al., 1986, Nature. 689-693). Exp7 is most similar to MgtB from S. typhimurium, which is one of three genetic loci responsible for the transport of Mg2+ (Snavely et al., 1991, J. Biol Chem. 266:815-823). The identified region contains the highly conserved aspartyl residue, which is the site for ATP dependent autophosphorylation. Based on the similarity to MgtB, the fusion in Exp7 probably occurred in the C-terminal region of the protein. A predicted model for the transmembrane loops of MgtB suggested that this region would be on the cytoplasmic surface (Snavely et al., 1991, J. Biol. Chem. 266:815-823). The data with the PhoA fusion to Exp7 suggests that location of this region on the cytoplasmic surface is not the case in S. pneumoniae.

Exp8 shows similarity to the family of traffic ATPases, alternatively called the ATP binding cassette (ABC) superfamily of transporters, which are found in both prokaryotes and eukaryotes (reviewed in Ames and Lecar, 1992, Faseb J. 6:2660-6) 15 (Figure 5H). Exp8 is most similar to the transmembrane proteins responsible for the translocation of bacterial RTX proteins such as the α-hemolysins, which are eukaryotic cytotoxins found in both Gram negative and Gram positive organisms (reviewed in Welch, 1991, Mol. Microbiol. 5:521-528). The fusion protein containing Exp8 is most similar to CyaB a component of the cya operon in Bordetella 20 pertussis (Glaser et al., 1988, Mol. Microbiol. 2:19-30; Glaser et al., 1988, EMBO J. 7:3997-4004). This locus produces the adenylate cyclase toxin which is a also member of the RTX family of bacterial toxins. It does not go without notice that the comA locus in S. pneumoniae is also a member of this family (Hui and Morrison, 1991. J. Bacteriol. 173:372-81).

25

The derived sequence for exp9 from two regions of the recovered insert are presented in Figure 4. Analysis of this sequence revealed that Exp9 is a member of the D-E-A-D protein family of ATP-dependent RNA helicases (for a review see (Schmid and Linder, 1992, Mol. Microbiol. 6:282-292)). It is most similar to DEAD from E. coli 30 (Figure 51) (Toone et al., 1991, J. Bacteriol. 173:3291-3302). A large number of helicases have been identified from many different organisms. At least five different

homologs have been identified in *E. coli* (Kalman et al., 1991, The New Biologist 3:886-895). The hallmark of these proteins is the conserved DEAD sequence within the B motif of an ATP binding domain (Walker et al., 1982, EMBO J. 1:945-951). The DEAD sequence was identified in the derived sequence from the 5' end of the 5 insert from exp9.

Two studies have suggested that different homologs in E. coli may play a role in translation by affecting ribosome assembly (Nishi et al., 1988, Nature. 336:496-498; Toone et al., 1991, J. Bacteriol. 173:3291-3302). No published studies have reported either export or membrane association of these proteins. Therefore it was surprising to identify a PhoA<sup>+</sup> mutant harboring this fusion. Subcellular fractionation clearly shows the majority of the fusion protein associated with the membrane fraction of the bacteria (Figure 6), although this could be an anomaly observed only with the fusion protein.

15

Recently, comF in B. subtilis has been shown to contain a similar RNA/DNA helicase with a DEAD sequence (Londonó - Vallejo and Dubnau, Mol. Microbiol.). Mutations in this locus render the bacteria transformation deficient. Subsequent studies have shown the helicase to be a membrane associated protein and it has been suggested that it may play a role in the transport of DNA during transformation (D. Dubnau, personal communication). Preliminary experiments have not shown a great difference in the transformability of a mutant expressing the Exp9-PhoA fusion. If there are a class of helicases associated with the membrane, it is tempting to speculate that Exp9 may be involved in the translation of polypeptides destined to be exported.

25

In conclusion, this Example demonstrates the development of a technique that successfully mutated and identified several genetic loci in *S. pneumoniae* that encode homologs of known exported proteins. It is clear from our results that the majority of the loci that have been identified encode exported proteins that play a role in several diverse processes that occur either at the cytoplasmic membrane or outside the bacteria. As with the use of PhoA mutagenesis in other organisms, a note of caution

- 63 -

is also advised with this technique in *S. pneumoniae*. Not all loci identified may encode exported proteins. It is certainly possible that due to several factors such as cell lysis some false positives may be generated. As demonstrated in the following Example, additional assays to demonstrate the functional activity of the mutant putative exported protein can be performed.

Given these results, the majority of the loci identified to date encode exported proteins, some of which play a role in signal transduction, protein translocation, cell wall biosynthesis, nutrient acquisition or maintaining a chemiosmotic balance.

10

# EXAMPLE 2: MUTATION OF SOME EXPORTED PROTEINS AFFECTS ADHERENCE

In this Example, the ability of encapsulated and unencapsulated pneumococci to

adhere to lung cells was determined. The results indicate that both types of
pneumococci adhere to mixed lung cells and to Type II lung cells, although the
preference was for type II cells. Also, the results suggest that the type 2 encapsulated
strain has a slightly greater ability to adhere than the unencapsulated variant.

20 The effect of mutations to exported proteins on the ability of the mutated S. pneumoniae strains to adhere to human umbilical vein endothelial cells (HUVEC) and lung Type II cells was also assayed. The results demonstrated that some of the exported proteins have direct or indirect roles in adhesion of S. pneumoniae to either HUVEC or lune cells. or both.

25

# Materials and Methods

Preparation of mixed and type II alveolar cells from rabbit.

As described by Dobbs and Mason (1979, J. Clin. Invest. 63:378-387), lungs were

removed from the rabbit, minced and digested with collagenase, elastase and DNase
for 60 min at 37°C. Large pieces were removed over a gauze filter and cells were

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pelleted and washed twice. The mixed lung cells were resuspended in 20 ml of calcium containing buffer supplemented with 0.5% albumin at a density of 10° per ml. Alveolar type II cells were purified from the mixed lung cell suspension by layering the suspension on an albumin gradient of 10 ml at 16.5 g% over 10 ml at 35 g% and centrifuged at 1200 rpm for 20 min at 4°C. The top 26 ml of the gradient were discarded and cells in the next 12 ml were harvested, washed and adjusted to a concentration of 10° cells per ml. Viability of the cells was greater than 90% by as assessed by Trypan blue exclusion, and greater than 80% of the cells contained osmiophilic lamellar bodies typical of Type II cells when examined by 0 electron microscopy.

# Adherence assay with mixed and Type II alveolar cells,

About 10<sup>3</sup> to 10<sup>5</sup> type II (encapsulated) or R6 (unencapsulated) pneumococci were added to 10<sup>4</sup> lung cells in a 1 ml volume for 30 min at 37 °C. Lung cells were separated from non-adherent bacteria by 6 rounds of washing by centrifugation at 270 x g for 5 min. Bacteria adherent to the final cell pellet were enumerated by plating and by Gram stain.

# HUVEC and Type II lung alveolar cell adherence assays.

- 20 HUVEC (Clonetics, San Diego, California) and Type II alveolar cell line cells (ATCC accession number A549) were cultured 4-8 days and then were transferred to Terasaki dishes 24 hours before the adherence assay was performed to allow formation of a confluent monolayer (Geelen et al., 1993, Infect. Immun. 61:1538-1543). Bacteria were labelled with fluorescein (Geelen et al., supra), and adjusted to a concentration of 5 x 10<sup>7</sup>, or to concentrations of 10<sup>3</sup>, 10<sup>4</sup> and 10<sup>7</sup> cfu per ml, and added in a volume of 5 μl to at least 6 wells. After incubation at 37 °C for 30 min, the plates were washed and fixed with PBS/glutaraldehyde 2.5%. Attached bacteria were enumerated visually using a Nikon Diaphot Inverted Microscope equipped with epifluorescence.
- 30

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An additional mutant strain of R6, SPRU25, was generated as described in Example 1. above.

# Results and Discussion

5

Adherence of encapsulated type 2 and unencapsulated R6 pneumococci to mixed lung cells (data not shown) was consistently 1-2 logs less at each inoculum than to purified Type II cells. This indicated that Type II cells were the preferred target for the bacteria. The concentration curve for Type II cells is shown in Figure 7. A consistent but statistically insignificant difference was noted between encapsulated an unencapsulated strains suggesting the type II strain might have a slightly greater ability to adhere than the unencapsulated variant.

Mutant strains (Table 1) were tested for the ability to adhere to HUVEC and lung

Type II cells. Strains SPRU98, SPRU42, SPRU40, SPRU25 and SPRU121

were found to have reduced adhesion activity compared to the R6 wildtype strain.

The adherence of other strains was not significantly affected by the mutation of exported proteins (data not shown).

- 20 The bacteria were titrated to 10<sup>3</sup>, 10<sup>6</sup> and 10<sup>7</sup> cfu per ml and tested for the ability to adhere to HUVEC (Figure 8) and lung Type II (Figure 9) cells. At the lowest concentration, the numbers of adherent bacteria were relatively the same between the adherence deficient mutants and R6. At 10<sup>6</sup>, and more notably at 10<sup>7</sup>, cfu per ml, the difference between binding by the mutants to both HUVEC and lung Type II cells
- 25 varied from significant to dramatic.

Homologies of the exported proteins of strains SPRU98, SPRU42, and SPRU40 are discussed in Example 1, above. SPRU121 represents a mutation of the amiA locus. The results of this experiment provide unexpected evidence that the AmiA exported protein is involved in adhesion. SPRU25 is a strain generated as described in Example 1, with a mutation at the exp10. No genes or proteins with homology to the

nucleic acid [SEQ ID NO:21] or amino acid [SEQ ID NO:22] sequences of this exported protein were found. The identified portion of the exp10 nucleotide and Exp10 amino acid sequences are shown in Figure 10.

5 These results clearly indicate that exported proteins of S. pneumoniae that play a role in adhesion of the bacterium to cells can be identified.

# EXAMPLE 3: PEPTIDE PERMEASES MODULATE TRANSFORMATION

- 10 The present example relates to further elucidation of the sequence and function of Expl, a mutant that consistently transformed 10 fold less than the parent strain. The complete sequence analysis and reconstitution of the altered locus revealed a gene, renamed plpA (permease like protein), which encodes a putative substrate binding protein belonging to the family of bacterial permeases responsible for pentide 15 transport. The derived amino acid sequence for this gene was 80% similar to AmiA. a peptide binding protein homolog from pneumococcus, and 50% similar over 230 amino acids to Spo0KA which is a regulatory element in the process of transformation and sporulation in Bacillus subtilis. PlpA fusions to alkaline phosphatase (PhoA) were shown to be membrane associated and labeled with [3H] palmitic acid which 20 probably serves as a membrane anchor. Experiments designed to define the roles of the plpA and ami determinants in the process of transformation showed that: 1] Mutants with defects in plpA were > 90% transformation deficient while ami mutants exhibited up to a four fold increase in transformation efficiency. 21 Compared to the parental strain, the onset of competence in an ami mutant occurred earlier in 25 logarithmic growth, while the onset was delayed in a plpA mutant. 31 The plpA mutation decreases the expression of a competence regulated locus. Since the permease mutants would fail to bind specific ligands, it seems likely that the substrate-permease interaction modulates the process of transformation.
- 30 This example demonstrates through mutational analysis that these two peptide permeases have distinct effects on the induction of competence as well as on

transformation efficiency. Therefore, we propose that peptide permeases mediate the process of transformation in pneumococcus through substrate binding and subsequent transport or signaling and that these substrates may be involved in the regulation of competence.

5

# Materials and Methods

Strains and Media. The strains of S. pneumoniae used in this Example are described in Example 1, in particular in Table 1. Table 5 lists other pneumococcal strains used in this study and summarizes their relevant characteristics. Escherichia coli strains used are described in Example 1.

Table 5. Bacterial strains of Streptococcus pneumoniae used in this study.

15	Strain	Relevant Characteristics	Integrated plasmid	Source
	R6x	hex', Parent strain	none	(Tiraby and Fox, 1973)
	SPRU58	plpA-phoA fusion	pHplp10	Current study
	SPRU98	plpA-phoA fusion	pHplp1	(Example 1)
	SPRU107	plpA*	pJplp1	Current study
20	SPRU114	amiA-	pJamiA1	Current study
	SPRU121	amiA-phoA fusion	pHamiA1	(Example 1)
	SPRU122	plpA <sup>-</sup>	pJplp9	Current study
	SPRU148	amiC	pJamiC1	Current study
	SPRU100	exp10-phoA fusion		manuscript in preparation
25	SPRU156	plpA , exp10-phoA fusion	pWplp9	manuscript in preparation

S. pneumoniae plating and culture conditions are described in Example 1. For labeling studies cultures were grown in a chemically defined media (C<sub>DEN</sub>) prepared as described elsewhere (Tomasz, 1964, Bacteriol. Proc. 64:29). E. coli were grown in either liquid Luria-Bertani media or on solid TSA media supplemented with 500 μg / ml erythromycin or 100 μg / ml ampicillin where appropriate. For the selection and maintenance of pneumococcus containing

chromosomally integrated plasmids, bacteria were grown in the presence of  $0.5~\mu g$  / ml erythromycin.

PhoA\* libraries and mutagenesis. Libraries of pneumococcal mutants expressing
5 PhoA fusions were created by insertional inactivation with the non replicating pneumococcal E. coli shuttle vectors pHRM100 or pHRM104. The pneumococcal E. coli shuttle vector pIDC9 was used for gene inactivation without the generation of phoA fusions. The plasmid constructs used for mutagenesis are shown in Fig.
7. The details for these procedures are described in Example 1.

10

Pneumococcal transformation. To screen large numbers of mutants for a decrease in transformation efficiency, single colonies were transferred to 96 well microtiter plates containing 250 μl of liquid media and chromosomal DNA (final concentration 1 μg / ml) from a streptomycin resistant strain of pneumococcus 15 (Str DNA). After incubation for 16 h at 37°C, 5 μl samples were plated onto solid media with and without antibiotic to determine transformation efficiency. Control strains produced approximately 10<sup>3</sup> Str transformants / ml while transformation deficient candidates produced less than 10<sup>4</sup> Str transformants / ml.

- 20 The permease mutants were assessed in a more defined transformation assay (Fig. 15). Stock cultures of bacteria were diluted to a cell density of approximately 10° cfu / ml in C+Y media containing Str DNA. This solution was dispensed into 250 μl aliquots in a 96 well microtiter plate and the bacteria were grown for 5 hours at 37°C to an OD<sub>600</sub> of approximately 0.6. Total bacteria and Str transformants were determined by serial dilution of the cultures onto solid media with and without antibiotic. Transformation efficiency was calculated as the percent of Str transformants / total number of bacteria and compared to the parent strain, R6x.
- 30 Competence profiles which assess transformation were generated from cultures grown in liquid media. Stocks of bacteria were diluted to a cell density of

approximately 10° cfu / ml into fresh C+Y media (10 ml) and grown at 37°C. Samples (500 μl) were withdrawn at timed intervals, frozen and stored in 10% glycerol at -70°C. These samples were thawed on ice then incubated with Str<sup>\*</sup> DNA for 30 min at 30°C. DNAse was added to a final concentration of 10 μg / ml to stop further DNA uptake and the cultures were transferred to 37°C for an additional 1.5 h to allow the expression of antibiotic resistance. Transformation efficiency was calculated as described above.

Recombinant DNA techniques. Standard DNA techniques including plasmid mini preparations, restriction endonuclease digests, ligations, transformation into E. coli and gel electrophoresis were according to standard protocols (Sambrook et al., 1989, supra). Restriction fragments used in cloning experiments were isolated from agarose gels using glass beads (Bio 101) or phenol extractions. Large scale plasmid preparations were prepared using the affinity columns according to the manufacturer's instructions (Qiagen).

Double stranded DNA sequencing was performed by the Sanger method (Sanger et al., 1977, Proc. Natl. Acad. Sci. USA 74:5463-67) using [a-35S]-dATP (New England Nuclear) and the Sequenase Version 2.0 kit (United States Biochemical Corp.), according to the manufacturer's instructions. Dimethysulphoxide (1% v/v) was added to the annealing and extension steps.

The polymerase chain reaction (PCR) was performed using the Gene Amp Kit (Perkin Elmer Cetus). Oligonucleotides were synthesized by Oligos Etc. Inc. or at the Protein Sequencing Facility at The Rockefeller University.

In vivo labeling of PlpA-PhoA. Frozen stocks of pneumococcus were resuspended in 4 ml of fresh  $C_{DEN}$  media and grown to an  $OD_{670}$  of 0.35 at 37°C. Each culture was supplemented with 100  $\mu$ Ci of [9,10-3H] palmitic acid (New England Nuclear) and grown for an additional 30 min. Cells were harvested by centrifugation and washed three times in phosphate buffered saline (PBS). The final cell pellet was

resuspended in 50  $\mu$ l of lysis buffer (PBS; DNAse, 10  $\mu$ g/ml; RNAse 10  $\mu$ g/ml; 5% [v/v] deoxycholate) and incubated for 10 min at 37°C. To immuno precipitate the PlpA-PhoA fusion protein the cell lysate was incubated with 20  $\mu$ l of anti-PhoA antibodies conjugated to Sepharose (5'3' Inc.) for 1 h at 4°C. The suspension was washed three times with equal volumes of PBS and once with 100  $\mu$ l 50 mM Tris-HCl pH 7.8, 0.5 mM dipotassium ethylenediaminetetra-acetate (EDTA). The final supernatant was discarded and the resin was resuspended in 30  $\mu$ l of SDS sample buffer, boiled for 5 min and subjected to SDS polyacrylamide gel electrophoresis and autoradiography.

10

Subcellular fractionation. Pneumococci were fractionated into subcellular components by a previously described technique (Hakenbeck et al., 1986, Antimicrob. Agents Chemother. 30:553-8). Briefly, bacteria were grown in 400 ml of C+Y medium to an OD<sub>600</sub> of 0.6 and isolated by centrifugation at 17,000 g 15 for 10 min. The cell pellet was resuspended in a total volume of 2 ml of TEPI (25 mM Tris-HCl, pH 8.0, 1 mM EDTA, 1 mM phenyl methyl sulfonyl fluoride, 20 µg/ml leupeptin and 20 µg / ml aprotinin). One half volume of washed glass beads was added and the mixture was vortexed for 15 to 20 min at 4°C until the cells were broken as documented by microscopic inspection. The suspension was 20 separated from the glass beads by filtration over a cintered glass funnel. The beads were washed with an additional 5 ml of TEPI. The combined solutions were centrifuged for 5 min at 500 g to separate cellular debris from cell wall material, bacterial membranes and the cytoplasmic contents. The supernatant was then spun for 15 min at 29,000 g. The pellet contained the cell wall fraction 25 while the supernatant was subjected to another centrifugation for 2 h at 370,000 g. The supernatant from this procedure contained the cytoplasmic fraction while the pellet contained the bacterial membranes. Samples from each fraction were evaluated for protein content and solubilized in SDS sample buffer for subsequent gel electrophoresis. PlpA-PhoA fusion proteins were detected with anti PhoA antiserum (5'3' Inc.) and visualized indirectly by enhanced chemiluminescence as described in Example 1.

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Recovery and sequencing of plpA. Fig. 18 shows a restriction endonuclease map of plpA and fragments of various subclones. Plasmids with fragments cloned into pHRM104 have the prefix H while those cloned into pJDC9 have the prefix J. The integrated plasmids pHplp1 and pHplp10 were isolated from SPRU98 and SPRU58 respectively by transformation into E. coli of spontaneously excised plasmids which contaminate chromosomal preparations of DNA. "Chromosome walking" was used to isolate most of plpA and the downstream region. The 500 bp insert from pHplp1 was cloned via KpnI into pJDC9 to produce pJplp1 which was shuttled back into pneumococcus to produce SPRU107. Chromosomal DNA from SPRU107 was digested with various restriction endonucleases that cut the vector once but not within the original fragment. The DNA was religated and transformed into E. coli with selection for the vector. Using this procedure PstI produced pJplp2 and HindIII produced pJplp3 which both extended the 3' region of the original fragment in pJplp1 by 190 bp, while SphI produced pJplp4 which 15 contained an additional 3.8 kb. Subcloning of a 900 bp internal fragment of pJplp4 into pJDC9 gave plasmid pJplp5, containing 630 bp downstream from the 3' end of plpA. A further 450 bp was isolated upstream from the original fragment using EcoRI (pJplp6). A 730 bp internal fragment of pJplp6 was cloned into pJDC9 giving pJplp7, and a 200 bp EcoRI/PstI internal fragment of pJplp6 was cloned into the appropriate sites of pJDC9 to produce pJplp8.

The region upstream of the original fragment of plpA was obtained by "homology cloning" using degenerate and specific oligonucleotides with chromosomal DNA in a polymerase chain reaction (PCR). The degenerate oligonucleotide, lipo1, (GCC GGA TCC GGW GTW CTT GCW GCW TGC where W is A + T) (SEQ ID NO: 49) was based on the lipoprotein precursor consensus motif present in AmiA (Alloing et al., 1990, Mol. Microbiol. 4:633-44) and SarA, a peptide permease binding protien homolog from S. gordonii (Jenkinson, 1992, Infect. Immun. 60:1225-8). The specific oligonucleotide, P1, (TAC AAG AGA CTA CTT GGA TCC) (SEQ ID NO: 50) was complimentary to the 5' end of the insert in plplp6. To prevent amplification of the highly homologous amiA gene, chromosomal DNA

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was used from SPRU114, which has a disrupted amiA. The chromosomal DNA was first digested with Xhol to give shorter templates. PCR conditions were 40 cycles at 94°C for 30 seconds for denaturing, 40°C for 30 seconds for annealing and 72°C for 1 min for extension. A 600 bp product was obtained, gel purified, 5 digested with BamHI and cloned into Bluescript KS (Stratagene) giving pBSplp9. The BamHI digested fragment was then subcloned into pJDC9 to produce pJplp9. This plasmid was transformed into pneumococcus to give SPRU122.

Generation of a plpA mutant containing a competence regulated gene fused to
alkaline phosphatase. The 600 bp BamHI fragment from pBSplp9 was ligated to
Saulila digested pWG5 (Lacks et al., 1991, gENE 104:11-17) resulting in
pWplp9. This plasmid was transformed into SPRU100, which contains a gene,
exp10, from the competence regulated rec locus, fused to phoA, giving SPRU156.
Correct integration of the vector into the chromosome was confirmed by PCR.

Alkaline phosphatase activity was measured as described in Example 1, but with a
final substrate concentration (p-nitrophenyl phosphate, Sigma) of 2.5 mg / ml.

The activity units were calculated using the following formula:

OD420 - 1.75 x OD550

20 time (h) x OD<sub>600</sub> (of resuspended culture)

Generation of ami mutants. Internal fragments of ami obtained by PCR and restriction endonuclease digestion were ligated into the appropriate shuttle vectors and transformed into pneumococcus to produce the various ami mutants.

25 Construction of the gene fusion between amiA and phoA has been previously described in Example 1 to give SPRU121. To obtain a truncated amiA, oligonucleotides ami1 (ACC GGA TCC TGC CAA CAA GCC TAA ATA TTC) (SEQ ID NO: 51) and ami2 (TTT GGA TCC GTT GGT TTA GCA AAA TCG CTT) (SEQ ID NO: 52) were used to generate a 720 bp product at the 5' end of amiA. This fragment was digested with HindIII and EcoRI, which are within the coding region of amiA, and the corresponding 500 bp fragment was cloned into

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pJDC9. The resulting plasmid pJamiA was transformed into pneumococcus to produce SPRU114. To inactivate amiC, oligonucleotides amiC1 (CTA TAC CTT GGT TCC TCG) (SEQ ID NO: 53) and amiC2 (TTT GGA TTC GGA ATT TCA CGA GTA GC) (SEQ ID NO: 54), which are internal to amiC, were used to generate a 300 bp product using PCR. The resulting fragment was digested with BamHI and cloned into pJDC9 producing the plasmid, pJamiC1, which was transformed into pneumococcus to produce SPRU148.

Northern analysis. RNA was prepared according to procedures adapted from

Simpson et al. (1993, FEMS Microbiol. Lett. 108:93-98). Bacteria were grown to
an OD<sub>620</sub> of 0.2 in C+Y media, pH 8.0. After centrifugation (12,000 g, 15 min,
4°C) the cell pellet was resuspended in 1/40 volume of lysing buffer (0.1%
deoxycholate, 8% sucrose, 70 mM dithiothreitol). SDS was added to 0.1% and
the suspension incubated at 37°C for 10 min. Cellular debris was removed and an
15 equal volume of cold 4 M lithium chloride was added to the supernatant. The
mixed suspension was left on ice overnight then centrifuged at 18,500 g, for 30
min at 4°C. The pellet containing RNA was resuspended in 1.2 ml cold sodium
acetate (100 mM, pH 7.0) and 0.5% SDS, extracted three times with an equal
volume of chloroform/isoamyl alcohol (25:24:1) and once with an equal
20 volume of chloroform/isoamyl alcohol (24:1). The RNA was precipitated with
ethanol and resuspended in sterile water. The yield and purity was determined by
spectrophotometry with a typical yield of 300 μg RNA from 80 ml of culture.

Samples of RNA were separated by electrophoresis in 1.2% agarose / 6.6% formaldehyde gels (Rosen and Villa-Komaroff, 1990, Focus 12:23-24). The gel was rinsed in water, and the RNA transferred to nitrocellulose filters (Schleicher and Schuell) by capillary blotting (Sambrook et al., 1989, supra).

Prehybridization was for 4 h in 0.2% Denhardts (1 x Denhardts is 1% Ficoll, 1% polyvinyl-pyrrolidone, 1% bovine serum albumin), 0.1% SDS, 3 x SSC (1 x SSC is 150 mM NaCl, 15 mM sodium citrate), 10 mM HEPES, 18 µg / ml denatured salmon sperm DNA and 10 µg / ml yeast tRNA at 65°C with gentle agitation.

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The DNA probe used to detect plpA transcripts was a 480 bp HindIII - BamHI fragment from pJplp9. For detection of amiA transcripts, the DNA probe was a 720 bp PCR product generated with oligonucleotides ami1 and ami2 (described above). The DNA fragments were labeled with [a-32P]-dCTP using the Nick 5 Translation System (New England Nuclear). Hybridization was at 65°C overnight. Hybridization washes were 2 x SSC, 0.5% SDS for 30 min at room temperature, followed by 3 x 30 min washes at 65°C in 1x SSC, 0.5 x SSC and 0.2 x SSC, all containing 0.5% SDS.

Results

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permease. To identify exported proteins in mutants as described in Example 1, supra, that participate in the process of transformation, 30 PhoA<sup>+</sup> mutants were

15 assessed for a decrease in transformation efficiency. In an assay designed to screen large numbers of mutants, transformation of a chromosomal mutation for streptomycin resistance (Str<sup>0</sup>) into the parental strain (R6x) produced approximately 10<sup>5</sup> cfu / ml Str<sup>4</sup> transformants. The PhoA<sup>+</sup> mutant, SPRU98 consistently showed a 90% reduction in the number of Str<sup>4</sup> transformants (10<sup>6</sup> cfu / ml).

20 Transformation of the PhoA<sup>+</sup> mutation the parent R6x produced strains that were both PhoA<sup>+</sup> and transformation deficient demonstrating that the mutation council by the capacity in page fixing week lighted to the defect in transformation. The growth

Identification of a transformation deficient mutant with a defect in a peptide

caused by the gene fusion was linked to the defect in transformation. The growth rate of SPRU98 was identical to the parental strain suggesting that the transformation deficient phenotype was not due to a pliotropic effect related to the growth of the organism (data not shown). Recovery and identification of the mutated locus in SPRU98 revealed plpA (permease like protein) (Fig. 11, SEQ ID NO:46), which corresponds to exp1. The derived amino acid sequence of plpA (SEQ ID NO: 47) Showed extensive similarity to the substrate binding proteins associated with bacterial permeases (for a review, see Tam and Saier, 1993, Microbiol. Rev. 57:320-346) with the greatest similarity to AmiA (60% sequence

identity) (Fig. 12A; SEQ ID NO: 48). Alignment of PlpA with the binding

proteins from the family of bacterial peptide permeases revealed several blocks of sequence similarity that suggest functional motifs common to all members of this family (Fig. 12B).

- 5 Most examples of peptide permeases have a genetic structure that consists of five genes that encode an exported substrate binding protein, and two integral membrane proteins and two membrane associated proteins that are responsible for substrate transport across the cytoplasmic membrane (for reviews, see Higgins, 1992, Annu. Rev. Cell. Biol. 8:67-113; Tam and Saier, 1993, supra). Sequence 10 analysis 630 bp immediately downstream and in the region 3.3 kb downstream of plpA, did not reveal any coding sequences that are homologs of these transport elements (data not shown). Therefore, if PlpA is coupled to substrate transport. then it may occur through the products of a distinct allele. This is not without precedence. In Salmonella typhimurium, the his I and argT genes encode the highly similar periplasmic binding proteins J and LAO. Both of these proteins deliver their substrates to the same membrane associated components (Higgins and Ames, 1981, Proc. Natl. Acad. Sci. USA 78:6038-42). Likewise, the periplasmic binding proteins LS-BP and LIV-BP of Escherichia coli, which transport leucine and branched chain amino acids, also utilize the same set of membrane-bound components (Landick and Oxender, 1985, J. Biol. Chem. 260:8257-61).
- We were unable to recover the 5' end of plpA perhaps due to toxicity of the expressed protein in E. coli. Similar difficulties have been encountered in cloning the genes of other pneumococcal permeases such as amiA and malX (Alloing et al., 1989, supra; Martin et al., 1989, Gene 80:227-238). Based on sequence similarity between the derived sequences of plpA and amiA all but 51 bp of the 5' end of the gene was cloned.

Membrane localization and post translational covalent modification of PlpA. Both

30 PlpA and AmiA contain the LYZCyz (Y= A, S, V, Q, T: Z= G, A: y= S, T,
G, A, N, Q, D, F: z = S, A, N, Q, G, W, E) consensus sequence in the N

terminus which is the signature motif for post translational lipid modification of lipoproteins in bacteria (Gilson et al., 1988, EMBO J. 7:3971-74; Yamaguchi et al., 1988, Cell 53:423-32). In gram positive organisms this modification serves to anchor these polypeptides to the cytoplasmic membrane (Gilson et al., 1988, supra). Specific examples of permease substrate binding proteins containing this consensus sequence include SarA from Streptococcus gordonii (Jenkinson, 1992, Infect. Immun. 60:1225-8), Spo0KA from B. subtilis (Perego et al., 1991, Mol. Micribiol. 5:173-185; Rudner et al., 1991, J. Bacteriol. 173:1388-98), TraC and PrgZ from E. faecalis (Ruhfel et al., 1993, J. Bacteriol. 175:5253-59; Tanimoto et al., 1993, J. Bacteriol 175:5260-64) and MalX from S. pneumoniae (Gilson et al., 1988, supra).

In support of this proposal, Fig. 13 shows that the PlpA-PhoA protein is exported and associated primarily with the cytoplasmic membranes. Small amounts were also detected in the cell wall fraction and in the culture supernatant suggesting that some of PlpA may be released from the membrane. This is also seen for the peptide binding protein OppA (Spo0KA) from B. subtilis, where OppA is initially associated with the cell but increasing proportions are released during growth (Perego et al., 1991, supra). Thus PlpA and OppA may be present on the outside of the cell in a releasable form as has been proposed for other lipoproteins in gram positive bacteria (Nielsen and Lampen, 1982, J. Bacteriol. 152:315-322). Although it cannot be ruled out that the presence of the fusion protein in these fractions does not reflect the location of the native molecule but rather the processing of a foreign protein, this seems unlikely, since other membrane associated PhoA fusions are firmly associated with cytoplasmic membranes.

Finally, a [3H] palmitic acid labeled 93 kDa protein corresponding to the PlpA-PhoA fusion protein was immuno precipitated from SPRU98 which contains a plpA-phoA genetic construct (Fig. 13, lower panel). In contrast, no similarly labeled protein was detected in either the parental control or in SRPU100 which contains an undefined PhoA fusion. This demonstrates in vivo post translational

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lipid modification of PlpA.

Transcriptional analysis of plpA and amiA. Transcripts of 2.2 kb were detected with probes specific for plpA and amiA in RNA preparations from R6x cells (Fig. 14). This is similar in size to the coding region for both genes. To eliminate the possibility of cross hybridization between the probes for plpA and amiA, high stringency washes were done after hybridization (see experimental procedures). The specificity of the probes was also demonstrated when RNA prepared from the mutant SPRU107, which contains a plasmid insertion in plpA, was probed with amiA and plpA. The amiA transcript remained at 2.2 kb while the plpA transcript shifted to 2.6 kb. In SPRU107, plpA is disrupted at bp 1474 by pJDC9. The plpA transcript would be 520 bp smaller than the full length transcript (1.7 kb), with an additional 800 bp from pJDC9 giving a transcript of about 2.5 kb, which is similar to the 2.6 kb transcript detected.

15

A single transcript corresponding to the size of plpA suggests that plpA is not part of an operon. This is confirmed by sequence analysis downstream of plpA which did not reveal any homologs to genes encoding transport elements commonly associated with peptide permeases (data not shown). Also, a potential rho independent transcription terminator was identified 21 bp downstream from the translational stop codon of plpA (Fig. 11).

Mutations in the PlpA and AmiA permeases have distinct effects on the process of transformation. To determine the effect of permeases during competence, we assessed the transformation efficiency of mutants with defects in either plpA or ami. In this assay, strains of bacteria were transformed with a selectable marker through a complete competence cycle followed by a subsequent outgrowth and then plated for the selection of the cells which have incorporated the antibiotic marker. Results are thus a measure of the total number of transformed cells during competence. Mutants that produced either truncated or PhoA fusions of PlpA exhibited a two to ten fold decrease in transformation efficiency (Fig. 15).

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In mutants with a disruption at Asp<sub>492</sub> of PlpA, the presence (SPRU98) or absence of PhoA (SPRU107), did not affect the 90% decrease in transformation efficiency. On the other hand, a mutant (SPRU122) producing a truncated PlpA at Asp<sub>192</sub> exhibited a 90% decrease in transformation efficiency, while in SPRU58 the fusion to PhoA at Leu<sub>197</sub> partially restored the parental phenotype. In this construct it is possible that PhoA conveys functionality by contributing to the chimera's tertiary structure thus affecting its ability to bind its substrate.

In contrast, mutants with defects in ami were transformation proficient. Mutants
that produced AmiA truncated at Propy either in the presence (SPRU121) or
absence (SPRU114) of PhoA showed a modest increase in transformation
efficiency (Fig. 15). Moreover, mutant SPRU148 with a disruption in AmiC
(Ile<sub>128</sub>) showed a four-fold increase in transformation efficiency. In this mutant we
presume that AmiA is produced and thus capable of binding its substrate.

15 Therefore, the increase observed with the amiC mutant suggests that substrate transport via the ami encoded transport complex may regulate transformation in addition to substrate binding by AmiA. Finally, even though PlpA and AmiA are highly related structures (60% sequence identity) the disparate effects observed with plpA and ami mutations on transformation efficiency suggest that substrate
20 specificity conveys these differences.

Transformation occurs during a single wave of competence early in logarithmic growth (Fig. 16). Therefore, regulation of this process may occur by either modifying the onset of competence (a shift in the curve) or by altering the expression of competence induced genes, leading to a change in the number of successfully transformed cells. To determine if the permeases regulate the process of transformation we compared the competence profiles of the permease mutants with the parental strain. This analysis measures the number of transformed cells in the population of cells at various stages of growth during a competence cycle.

30 Fig. 16 shows a single wave of competence for the parental strain (R6x) with a

maximal transformation efficiency of 0.26% at an OD of 0.12. This

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corresponds to a cell density of approximately 10' cfu / ml. A plpA mutant (SPRU107) underwent a similar wave of transformation with a maximal transformation efficiency of only 0.06% at a higher cell density. In contrast, an amiA mutant (SPRU114) underwent a wave of transformation that persisted over more than one doubling time with a maximal transformation efficiency of 0.75%. The onset of the competence cycle in SPRU114 occurred at an earlier cell density beginning by an OD<sub>620</sub> of 0.03. From this data we conclude that mutations in either permease has a dual effect on the process of transformation, affecting both the induction of the competence cycle as well as modulating the successful number 0 of transformants.

A mutation in plpA causes a decrease in the expression of a competence regulated locus. The rec locus in pneumococcus, which is required for genetic transformation, contains two genes, exp10 and recA. Results with a translational exp10 - phoA gene fusion have demonstrated a 10 fold increase in enzyme activity with the induction of competence demonstrating that this is a competence regulated locus. To determine if the peptide permeases directly affect the expression of this competence induced locus, we constructed a mutant (SPRU156) with a null mutation in plpA and the exp10 - phoA gene fusion. By measuring alkaline 20 phosphatase activity during growth, we showed that compared to an isogenic strain (SPRU100), the mutant harboring the plpA mutation demonstrated almost a two fold decrease in the expression of the exp10-phoA fusion (Fig. 17). Therefore, these results show that at least plpA directly affects the signaling cascade responsible for the expression of a competence regulated gene required for transformation.

### Discussion

The newly identified export protein Exp1, is encoded by the genetic determinant, renamed herein plpA. This locus, along with the ami locus, modulates the process of transformation in S. pneumoniae. Both loci encode highly similar peptide binding proteins (PlpA, AmiA) that are members of a growing family of bacterial

permeases responsible for the transport of small peptides (Fig. 12B). Examples of these peptide binding proteins have been associated with the process of genetic transfer in several bacteria. In B. subtilis, inactivation of spo0KA, the first gene of an operon with components homologous to the peptide permeases, caused a decrease in transformation efficiency as well as arresting sporulation (Perego et al., 1991, supra; Rudner et al., 1991, supra). The substrate for Spo0KA is not known. B. subtilis produces at least one extracellular differentiation factor that is required for sporulation (Grossman and Losick, 1988, supra) and it has been proposed that this transport system could be involved in sensing this extracellular 10 peptide factor which may be required for competence and sporulation.

Conjugal transfer of a number of plasmids in *E. faecalis* is controlled by small extracellular peptide pheromones. Recent genetic analyses have identified two plasmid encoded genes, prgZ and traC, whose derived products are homologous to the peptide binding proteins. Experimental evidence suggests that these proteins may bind the peptide pheromones thus mediating the signal that controls conjugation (Ruhfel et al., 1993, supra; Tanimoto et al., 1993, supra). The absence of membrane transport elements is a common feature between the prgZ, traC and plpA determinants which implies either that transport is not required for signal transduction or that a distinct allele is required for transport.

Mutations in plpA and ami cause a decrease or an increase in transformation efficiency, respectively. In addition, mutations in these loci affect the induction of the growth stage specific competent state. Compared to the parent strain, a mutation in ami induces an earlier onset of competence while a mutation in plpA delays this induction. Furthermore, a translational fusion to a competence regulated locus has shown that a mutation in plpA directly affects the expression of a gene required for the process of transformation. Given that the induction of competence occurs as a function of cell density (Tomasz, 1966, J. Bacteriol. 30 91:1050-61), it is reasonable to propose that these permeases serve as regulatory elements that modulate the cell density dependent induction of competence by

mediating the binding and or transport of signaling molecules. Small peptides which are the presumed substrates for permeases in other bacteria or the extracellular pneumococcal activator protein are likely candidates as ligands for these permeases. Because peptide permease defective mutants of Salmonella typhimurium and Escherichia coli fail to recycle cell wall peptides released into culture media, it has been proposed that these permeases bind and transport cell wall peptides (Goodell and Higgins, 1987, J. Bacteriol. 169:3861-65; Park, 1993, J. Bacteriol. 175:7-11). Thus, cell wall peptides are likely candidates. Recent genetic evidence suggests that divalent cation (Ni<sup>2+</sup>) transport is also coupled to peptide permease function in E. coli (Navarro et al., 1993, Mol. Microbiol. 9:1181-91). It has also been shown that extracellular Ca<sup>2+</sup> coupled to intracellular transport can affect transformation (Trombe, 1993, J. Gen. Microbiol. 139:433-439; Trombe et al., 1992, J. Gen. Microbiol. 138:77-84). Therefore, peptide permease mediated divalent cation transport is also a viable model for intracellular signaling and subsequent modulation of transformation.

## EXAMPLE 4:

## A PYRUVATE OXIDASE HOMOLOG REGULATES ADHERENCE

- 20 The present Example describes isolation and sequence determination of an Exp mutant that encodes a pyruvate oxidase homolog. This new protein regulates bacterial adherence to eucaryotic cells.
- Bacterial adhesion to epithelial cells of the nasopharymx is recognized as a

  25 requirement for colonization of the mucosal surface and infection. Pneumococcal
  cell wall and proteins of the bacterial surface mediate attachment to eukaryotic
  cells. The molecular determinants that pneumococcus recognizes on the surface of
  the eucaryotic cell are complex sugars, particularly GlcNAcβ1-3Gal or GalNAcβ14Gal carbohydrate moieties.

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Mutants, as described in Example 1, supra, were screened for loss of binding to

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type II lung cells (T2LC), human endothial cells (HUVEC), and to GlcNAcβ1-3Gal sugar receptors in a hemagglutination assay that reflects adherence to cells in the nasopharynx.

One out of 92 independent mutants, named Pad1 (gneumococcal adherence 1), exhibited an inability to hemagglutinate the GlcNAcβ1-3Gal sugar receptor on neuraminidase-treated bovine erythrocytes as described (Andersson et al., see Example 2). Subsequently, this mutant has been renamed PoxB.
 Hemagglutination of neuraminidase treated bovine erythrocytes reflects adherence
 to cells in the nasopharynx. Directed mutagenesis of the parent strain inactivating

padI reconfirmed that the loss of hemagglutination was linked to this locus.

This mutant also exhibited a greater than 70% decrease in adhesion to T2LCs and HUVECs, as shown in Figure 19.

Recovery and reconstitution of the mutated locus pad1 revealed an open reading frame of 1.8 kb with sequence similarity to enzymes in the acetohydroxy acid synthase-pyruvate oxidase family. In particular, pad1 shares 51% sequence similarity with recombinant pax, and 32% similarity with paxB. Targeted genetic disruption of the locus in the parent strain showed that mutation at this locus was responsible for the loss of adherence in all three assays.

Subcellular fractionation of a mutant that expressed a Padl-PhoA fusion showed that the protein localized to the membrane and the cytoplasm (Figure 20A).

25 Comparison of antigenic surface components in the parent and mutant strain showed that loss of a 17 kDa polypeptide that did not correspond to Pad1 (Figure 20B).

These results indicate that Pad1 affects pneumococcal adherence to multiple cell 30 types, possibly by regulating the expression of bacterial adhesins.

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The Pad1 mutant required acetate for growth in a chemically defined media (Figures 21 and 22). Growth in acetate restored the adhesion properties of the bacteria to both lung and endothelial cells.

5 The nucleotide sequence information for the pad1 promoter region shows a putative -35 site, a -10 taatat sequence, a ribosome binding site, and a translation start site (Figure 23) (SEQ ID NO: 55). The deduced protein translation of this region is also provided (Figure 23) (SEQ ID NO: 56).

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This invention may be embodied in other forms or carried out in other ways without departing from the spirit or essential characteristics thereof. The present disclosure is therefore to be considered as in all respects illustrative and not restrictive, the scope of the invention being indicated by the appended Claims, and all changes which come within the meaning and range of equivalency are intended to be embraced therein.

It is also to be understood that all base pair sizes given for nucleotides and all molecular weight information for proteins are approximate and are used for the 10 purpose of description.

 Various references are cited throughout this specification, each of which is incorporated herein by reference in its entirety.

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### SEQUENCE LISTING

## (1) GENERAL INFORMATION:

- (i) APPLICANT: Rockefeller University, The Masure Ph.D., H. Robert Pearce, Barbara J. Tuomanen, Elaine
- (ii) TITLE OF INVENTION: BACTERIAL EXPORTED PROTEINS AND ACELLULAR VACCINES BASED THEREON
- (iii) NUMBER OF SEQUENCES: 56
- (iv) CORRESPONDENCE ADDRESS:
- (A) ADDRESSEE: Klauber & Jackson
  - (B) STREET: 411 Hackensack Avenue
  - (C) CITY: Hackensack
  - (D) STATE: New Jersey (E) COUNTRY: USA
  - (F) ZIP: 07601
- (v) COMPUTER READABLE FORM:
- - (A) MEDIUM TYPE: Floppy disk
  - (B) COMPUTER: IBM PC compatible
    (C) OPERATING SYSTEM: PC-DOS/MS-DOS
  - (D) SOFTWARE: PatentIn Release #1.0, Version #1.25
- (vi) CURRENT APPLICATION DATA:
  - (A) APPLICATION NUMBER: WO to be assigned
  - (B) FILING DATE: 01-SEP-1994
  - (C) CLASSIFICATION:
- (vii) PRIOR APPLICATION DATA:
  - (A) APPLICATION NUMBER: US 08/245,511
  - (B) FILING DATE: 18-MAY-1994
- (vii) PRIOR APPLICATION DATA:
  - (A) APPLICATION NUMBER: US 08/116,541
    (B) FILING DATE: 01-SEP-1994
- (viii) ATTORNEY/AGENT INFORMATION:
  - (A) NAME: Jackson Esq., David A.
  - (B) REGISTRATION NUMBER: 26,742 (C) REFERENCE/DOCKET NUMBER: 600-1-069 PCT
  - (ix) TELECOMMUNICATION INFORMATION:
    - (A) TELEPHONE: 201 487-5800 (B) TELEFAX: 201 343-1684 (C) TELEX: 133521
- (2) INFORMATION FOR SEQ ID NO:1:
  - (i) SEQUENCE CHARACTERISTICS:
    - (A) LENGTH: 490 base pairs
    - (B) TYPE: nucleic acid (C) STRANDEDNESS: both
    - (D) TOPOLOGY: unknown -
  - (ii) MOLECULE TYPE: DNA (genomic)
  - (iii) HYPOTHETICAL: NO

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(vi) ORIGINAL SOURCE:	(iv)	ANTI-SENSE: NO
	(vi)	ORIGINAL SOURCE: (A) ORGANISM: Streptococcus pneumoni.

(A) ORGANISM: Streptococcus pneumoniae
(B) STRAIN: R6

(vii) IMMEDIATE SOURCE: (B) CLONE: SPRU98

(ix) FEATURE: (A) NAME/KEY: CDS (B) LOCATION: 1..490

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:1:

	 		 	 	-					
							ACT Thr			48
							GTT Val			96
							GTC Val 45			144
							GAT Asp			192
							AAA Lys			240
							ATG Met			288
							ATG Met			336
							GAT Asp 125			384
							GCT Ala			432
							TGG Trp			480
TTT Phe	GAT Asp	С								490

## (2) INFORMATION FOR SEQ ID NO:2:

(i) SEQUENCE CHARACTERISTICS:
(A) LENGTH: 163 amino acids

- 87 -

- (B) TYPE: amino acid
- (D) TOPOLOGY: linear
- (ii) MOLECULE TYPE: protein
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO:2:

Asp Arg Thr Ala Tyr Ala Ser Gln Leu Asn Gly Gln Thr Gly Ala Ser

Lys Ile Leu Arg Asn Leu Phe Val Pro Pro Thr Phe Val Gln Ala Asp

Gly Lys Asn Phe Gly Asp Met Val Lys Glu Lys Leu Val Thr Tyr Gly 40

Asp Glu Trp Lys Asp Val Asn Leu Ala Asp Ser Gln Asp Gly Leu Tyr

Asn Pro Glu Lys Ala Lys Ala Glu Phe Ala Lys Ala Lys Ser Ala Leu

Gln Ala Glu Gly Val Thr Phe Pro Ile His Leu Asp Met Pro Val Asp

Gln Thr Ala Thr Thr Lys Val Gln Arg Val Gln Ser Met Lys Gln Ser

Leu Glu Ala Thr Leu Gly Ala Asp Asn Val Ile Ile Asp Ile Gln Gln

Leu Gln Lys Asp Glu Val Asn Asn Ile Thr Tyr Phe Ala Glu Asn Ala

Ala Gly Glu Asp Trp Asp Leu Ser Asp Asn Val Gly Trp Gly Pro Asp

Phe Ala Asp

- (2) INFORMATION FOR SEO ID NO:3:
  - (i) SEQUENCE CHARACTERISTICS:
    - (A) LENGTH: 960 base pairs
    - (B) TYPE: nucleic acid
    - (C) STRANDEDNESS: both (D) TOPOLOGY: unknown
  - (ii) MOLECULE TYPE: DNA (genomic)
  - (iii) HYPOTHETICAL: NO

  - (iv) ANTI-SENSE: NO
  - (vi) ORIGINAL SOURCE:
    - (A) ORGANISM: Streptococcus pneumoniae
    - (B) STRAIN: R6
  - (vii) IMMEDIATE SOURCE: (B) CLONE: SPRU42
    - (ix) FEATURE:
- (A) NAME/KEY: CDS (B) LOCATION: 1.,960

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(xi) SEQUENCE DESCRIPTION: SEQ ID NO:3:

												CTC Leu				48
												GAT Asp				96
												CGC Arg 45				144
CAC His	AGG Arg 50	GGG Gly	ATT Ile	GAT Asp	ACC Thr	ATC Ile 55	CGT Arg	ATC Ile	CTG Leu	GGA Gly	GCT Ala 60	TTC Phe	TTG Leu	CGC Arg	AAT Asn	192
												ACT Thr				240
												CAG Gln				288
												GAA Glu				336
ACC Thr	AAG Lys	CAA Gln 115	GAA Glu	ATC Ile	TTG Leu	ACC Thr	TAC Tyr 120	тат Туг	ATA Ile	AAT Asn	AAG Lys	GTC Val 125	TAC Tyr	ATG Met	TCT Ser	384
												TAC Tyr				432
												CTG Leu				480
												CCA Pro				528
												AAT Asn				576
ATC Ile	TCT Ser	GCT Ala 195	GAA Glu	CAG G1n	TAT Tyr	GAG Glu	AAA Lys 200	GCA Ala	GTC Val	AAT Asn	ACA Thr	CCA Pro 205	ATT Ile	ACT Thr	GAT Asp	624
												GCT Ala				. 672
AAT Asn 225	TAC Tyr	CTC Leu	AAG Lys	GAA Glu	GTC Val 230	ATC Ile	AAT Asn	CAA Gln	GTT Val	GAA Glu 235	GAA Glu	GAA Glu	ACA Thr	GGC Gly	TAT Tyr 240	720
AAC Asn	CTA Leu	CTC Leu	ACA Thr	ACT Thr 245	GGG Gly	ATG Met	GAT Asp	GTC Val	TAC Tyr 250	ACA Thr	AAT Asn	GTA Val	GAC Asp	CAA Gln 255	GAA Glu	768

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						GAA Glu		816
						GTT Val 285		864
						CAG Gln		912
						CGC Arg		960

### (2) INFORMATION FOR SEQ ID NO:4:

- (i) SEQUENCE CHARACTERISTICS:
  - (A) LENGTH: 320 amino acids (B) TYPE: amino acid

  - (D) TOPOLOGY: linear
- (ii) MOLECULE TYPE: protein

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:4: Thr Thr Ser Ser Lys Ile Tyr Asp Asn Lys Asn Gln Leu Ile Ala Asp Leu Gly Ser Glu Arg Arg Val Asn Ala Gln Ala Asn Asp Ile Pro Thr Asp Leu Val Lys Ala Ile Val Ser Ile Glu Asp His Arg Phe Phe Asp His Arg Gly Ile Asp Thr Ile Arg Ile Leu Gly Ala Phe Leu Arg Asn 50 55 60 Leu Gln Ser Asn Ser Leu Gln Gly Gly Ser Ala Leu Thr Gln Gln Leu 65 70 75 80 Ile Lys Leu Thr Tyr Phe Ser Thr Ser Thr Ser Asp Gln Thr Ile Ser 85 90 95 Arg Lys Ala Gln Glu Ala Trp Leu Ala Ile Gln Leu Glu Gln Lys Ala Thr Lys Gln Glu Ile Leu Thr Tyr Tyr Ile Asn Lys Val Tyr Met Ser Asn Gly Asn Tyr Gly Met Gln Thr Ala Ala Gln Asn Tyr Tyr Gly Lys 130 135 140 Asp Leu Asn Asn Leu Ser Leu Pro Gln Leu Ala Leu Leu Ala Gly Met 145 150 155 160 Pro Gln Ala Pro Asn Gln Tyr Asp Pro Tyr Ser His Pro Glu Ala Ala 165 170 175 Gln Asp Arg Arg Asn Leu Val Leu Ser Glu Met Lys Asn Gln Gly Tyr. 180 185 190 Ile Ser Ala Glu Gln Tyr Glu Lys Ala Val Asn Thr Pro Ile Thr Asp

- 90 -

Asp Pro Leu Ser Ile Asn Gln Gln Gly Asn Asp Arg Gly Arg Gln Tyr 15 15 1 15 1 15 1 15 1 15 1 15 1 15 1			195					2.00					205					
Asn Leu Leu Thr Thr Gly Met Asp Val Tyr Thr Asn Val Asp Gln Glu 245  Ala Gln Lys His Leu Trp Asp Ile Tyr Asn Thr Asp Glu Tyr Val Ala 260  Tyr Pro Asp Asp Glu Leu Gln Val Ala Ser Thr Ile Val Asp Val Ser 275  Asn Gly Lys Val Ile Ala Gln Leu Gly Ala Arg His Gln Ser Ser Asn 290  Val Ser Phe Gly Ile Asn Gln Ala Val Glu Thr Asn Arg Asp Trp Gly 310  (2) INFORMATION FOR SEQ ID NO:5:  (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 520 base pairs (B) Tyrs Inucleic acid (C) STRANDEUNSES: both (D) TOPCLOGY: unknown  (ii) MOLECULE TYPE: DNA (genomic)  (iii) HYPOTHETICAL: NO  (iv) ANTI-SENSE: NO  (vi) ORIGINAL SOURCE: (A) ORGANISM: Streptococcus pneumoniae (B) STRAIN: R6  (vii) IMMEDIATE SOURCE: (B) CLONE: SPRU40  (ix) FEATURE: (A) NAMEK/KEY: CDS (B) LOCATION: 1519  (xi) SEQUENCE DESCRIPTION: SEQ ID NO:5:  GAT CCT CTA TCT ATC AAT CAA CAA GGG AAT GAC CGT GGT CGC CAA TAT Asp Pro Leu Ser Ile Asn Gln Gln Gly Asn Asp Arg Gly Arg Gln Tyr 1  CGA ACT GGG ATT TAT TAT CAG GAT GAA GCA GAT TTG CCA GCT ATC TAC Arg Thr Gly Ile Tyr Tyr Gln Asp Glu Ala Asp Leu Pro Ala Val  ACA GTG GTG CAG GAG CAG GAA CGC ATG CTG GGT CGA AAA ATT CAA CTG GGT GTG CGC GAA AAT TAT CAI GIG Glu Glu Any Met Leu Gly Arg Sile Ala Val  ACA GTG GTG CAG GAG CAG GAA CGC ATG CTG GGT CGA AAA ATT CAA CTG GGT GTG CGC AAA ATT CTC VAL CATG GTG GTG CAC GAA AAT GAA CAA GTG GTG CGC CAA AAA CTAA CT	Gly		Gln	Ser	Leu	ГЛЗ		Ala	Ser	Asn	Tyr		Ala	Tyr	Met	Asp		
Ala Gln Lys Leu Trp Asp Ile Tyr Asn Thr Asp Glu Tyr Val Ala 260  Tyr Pro Asp Asp Glu Leu Gln Val Ala Ser Thr Ile Val Asp Val Ser 285  Asn Gly Lys Val Ile Ala Gln Leu Gly Ala Arg His Gln Ser Ser Asn 290  Val Ser Phe Gly Ile Asn Gln Ala Val Glu Thr Asn Arg Asp Trp Gly 310  (2) INFORMATION FOR SEQ ID NO:5:  (1) SEQUENCE CHARACTERISTICS: (A) LENGTH: 520 base pairs (B) Type: nucleic acid (C) STRANDEDNESS: both (D) TOPOLOGY: unknown  (ii) MOLECULE TYPE: DNA (genomic)  (iii) Hypothetical: NO  (iv) ANTI-SENSE: NO  (vi) ORIGINAL SOURCE: (B) STRAIN: R6  (vii) IMMEDIATE SOURCE: (B) CLONE: SPRU40  (ix) FEARIN: R6  (vii) SEQUENCE DESCRIPTION: SEQ ID NO:5:  GAT CCT CTA TCT ATC AAT CAA CAA GGG AAT GAC CGT GGT CGC CAA TAT ASp Pro Leu Ser Ile Asn Gln Gln Sen Asp Arg Gly Arg Gln Tyr 1 1 15  CGA ACT GGG ATT TAT TAT CAG GAT GAA GCA GAT TTG CCA GCT ATC TAC ARG Thr Gly Ile Tyr Tyr Gln Asp Glu Ala Asp Leu Pro Ala Ile Tyr 20  ACA GTG GTG CAG GAG CAG GAA CGC ATG CTG GGT CGA GAA ATT CAC THR Val Val Gln Glu Gln Glu Arg Met Leu Gly Arg Leu Pro Ala Ile Tyr 10 10 10 10 10 10 10 10 10 10 10 10 10		Tyr	Leu	Lys	Glu		Ile	Asn	Gln	Val		Glu	Glu	Thr	Gly			
Tyr Pro Asp Asp Glu Leu Gln Val Ala Ser Thr Ile Val Asp Val Ser 275 285 285 285 285 285 285 285 285 285 28	Asn	Leu	Leu	Thr		Gly	Met	Asp	Val	Tyr 250	Thr	Asn	Val	Asp				
Asn Gly Lys Val Ile Ala Gln Leu Gly Ala Arg His Gln Ser Ser Asn 290  Val Ser Phe Gly Ile Asn Gln Ala Val Glu Thr Asn Arg Asp Trp Gly 310  (2) INFORMATION FOR SEQ ID NO:5:  (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 520 base pairs (B) Type: nucleic acid (C) STRANBEDNESS: both (D) TOPOLOGY: unknown  (ii) MOLECULE TYPE: DNA (genomic)  (iii) HYPOTHETICAL: NO  (vi) ANTI-SENSE: NO  (vi) ORIGINAL SOURCE: (A) ORGANISM: Streptococcus pneumoniae (B) STRAIN: RG  (vii) IMMEDIATE SOURCE: (A) ORGANISM: Streptococcus pneumoniae (ix) FEATURE: (A) ANAMEKEY: CDS (B) LOCATION: 1519  (xi) SEQUENCE DESCRIPTION: SEQ ID NO:5:  GAT CCT CTA TCT ATC AAT CAA CAA GGG AAT GAC CGT GGT CGC CAA TAT ASP Pro Leu Ser Ile Asn Gln Gly Asn Asp Arg Gly Arg Gln Tyr 1 1 15  CGA ACT GGG ATT TAT TAT CAG GAT GAA GCA GAT TTG CCA GCT ATC TAC ARG Thr Gly Ile Tyr Tyr Gln Asp Glu Ala Asp Leu Pro Ala Ile Tyr 20 25  ACA GTG GTG CAG GAG CAG GAA CGC ATG CTG GGT CGA AAA ATT CAA THY Val Val Gln Glu Gln Glu Ang Met Leu Gly Arg Sile Ala Val  46  ACA GTG GTG CAG GAG CAG GAA CGC ATG CTG GGT CGA AAA ATT GCA GTA Thr Val Val Gln Glu Gln Glu Ang Met Leu Gly Arg Sile Ala Val  46	Ala	Gln	Lys		Leu	Trp	Asp	Ile		Asn	Thr	Ąsp	Glu		Val	Ala		
Val Ser Phe Gly Ile Asn Gln Ala Val Glu Thr Asn Arg Asp Trp Gly 305  Val Ser Phe Gly Ile Asn Gln Ala Val Glu Thr Asn Arg Asp Trp Gly 305  (2) INFORMATION FOR SEQ ID NO:5:  (3) SEQUENCE CHARACTERISTICS: (A) LENGTH: \$20 base pairs (B) TrpE: nucleic acid (C) STRANDEDNESS: both (D) TOPOLOGY: unknown  (ii) MOLECULE TYPE: DNA (genomic)  (iii) HYPOTHETICAL: NO  (iv) ANTI-SENSE: NO  (vi) ORIGINAL SOURCE: (A) ORGANISM: Streptococcus pneumoniae (B) STRAIN: R6  (vii) IMMEDIATE SOURCE: (B) CLONE: SPRU40  (ix) FEATURE: (A) NAME/KEY: CDS (B) LOCATION: 1519  (xi) SEQUENCE DESCRIPTION: SEQ ID NO:5:  GAT CCT CTA TCT ATC AAT CAA CAG GGG AAT GAC CGT GGT CGC CAA TAT Asp Pro Leu Ser Ile Asn Gln Gln Gly Asn Asp Arg Gly Arg Gln Tyr 1  CGA ACT GGG ATT TAT TAT CAG GAT CAA GCA GAT TTG CCA GCT ATC TAC Arg Thr Gly Ile Tyr Gln Asp Glu Ala Asp Leu Pro Ala Ile Tyr 20  ACA GTG GTG CAG GAG CAG GAA CGC ATG CTG GGT CGA AAA THA THY Aly Val Val Gln Glu Gln Glu Arg Met Leu Gly Arg Lys Ile Ala Val  40	Tyr	Pro		Asp	Glu	Leu	Gln		Ala	Ser	Thr	Ile		Asp	Val	Ser		
(2) INFORMATION FOR SEQ ID NO:5:  (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 520 base pairs (B) TYPE INUCLEC seld (C) STRANDENNES: both (D) TOPOLOGY: unknown  (ii) MOLECULE TYPE: DNA (genomic)  (iii) HYPOTHETICAL: NO (iv) ANTI-SENSE: NO (vi) ORIGINAL SOURCE: (A) ORGANISM: Streptococcus pneumoniae (B) STRAIN: R6  (vii) IMMEDIATE SOURCE: (B) CLONE: SPRU40  (ix) FEATURE: (A) NAMEK/KEY: CDS (B) LOCATION: 1519  (xi) SEQUENCE DESCRIPTION: SEQ ID NO:5:  GAT CCT CTA TCT ATC AAT CAA CAA GGG AAT GAC CGT GGT CGC CAA TAT ABP PRO Leu Ser I1e Asn Gln Gln Gly Asn Asp Arg Gly Arg Gln Tyr 1  CGA ACT GGG ATT TAT TAT CAG GAT GAA GCA GAT TTG CCA GCT ATC TAC ARG Thr Gly 11e Tyr Tyr Gln Asp Glu Ala Asp Leu Pro Ala 11e Tyr 20  ACA GTG GTG CAG GAG CAG GAA CGC ATG CTG GGT CGA GAA ATT CTAC TAT VAI Val Val Gln Glu Gln Glu Ang Met Leu Gly Arg Sile Ala Val  46  ACA GTG GTG CAG GAG CAG GAA CGC ATG CTG GGT CGA GAA ATT CAC ATG TAT VAI Val Val Gln Glu Gln Glu Ang Met Leu Gly Arg Sile Ala Val  46  ACA GTG GTG CAG GAG CAG GAA CGC ATG CTG GGT CGA AAG ATT GCA GTA Thr Val Val Gln Glu Gln Glu Ang Met Leu Gly Arg Sile Ala Val			Lys	Val	Ile	Ala		Leu	Gly	Ala	Arg		Gln	Ser	Ser	Asn		
(i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 520 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: both (D) TOPOLOGY: unknown  (ii) MOLECULE TYPE: DNA (genomic)  (iii) HYPOTHETICAL: NO (iv) ANTI-SENSE: NO (vi) ORIGINAL SOURCE: (A) ORGANISM: Streptococcus pneumoniae (B) STRAIN: R6 (vi) IMMEDIATE SOURCE: (B) CLONE: SFRU40  (ix) FEATURE. (A) NAMEN/KEY: CDS (B) LOCATION: 1519  (xi) SEQUENCE DESCRIPTION: SEQ ID NO:5:  GAT CCT CTA TCT ATC AAT CAA CAA GGG AAT GAC CGT GGT CGC CAA TAT ABP PRO Leu Ser Ile Asn Gin Gly Asn Asp Arg Gly Arg Gin Tyr 1 1 5 10  CGA ACT GGG ATT TAT TAT CAG GAT GAA GCA GAT TTG CCA GCT ATC TAC Arg Thr Gly Ile Tyr Tyr Gin Asp Glu Ala Asp Leu Pro Ala Ile Tyr 20 25 30  ACA GTG GTG CAG GAG CAG GAA CGC ATG CTG GGT CGA GAA ATT GCA GTA Thr Val Val Gln Glu Gln Glu Amp Met Leu Gly Arg Sile Ala Val 35 40		Ser	Phe	Gly			Gln	Ala	Val	Glu		Asn	Arg	Asp	Trp			
(B) CLONE: SPRU40  (ix) FEATURE: (A) NAMEKEY: CDS (B) LOCATION: 1519  (xi) SEQUENCE DESCRIPTION: SEQ ID NO:5:  GAT CCT CTA TCT ATC AAT CAA CAA GGG AAT GAC CGT GGT CGC CAA TAT ASP PRO Leu Ser 11e Asn Gln Gln Gly Asn Asp Arg Gly Arg Gln Tyr 1 15 10 15 15  CGA ACT GGG ATT TAT TAT CAG GAT GAA GCA GAT TTG CCA GCT ATC TAC Arg Thr Gly 11e Tyr Tyr Gln Asp Glu Ala Asp Leu Pro Ala Ile Tyr 20  ACA GTG GTG CAG GAG CAG GAA CGC ATG CTG GGT CGA AAG ATT GCA GTA Thr Val Val Gln Glu Gln Glu Arg Met Leu Gly Arg Lys Ile Ala Val 315  40		(ii) (iii) (iii)	SEQ () (I) (I) (I) (I) (I) (I) (I) (I) (I)	QUENCA) LECUI COTHE	CE CIENGTI (PE: TRANI (POLO LE TI CTICA ENSE:	HARACH: 52 nuclocednicody: VPE: AL: NO DURCE	TTERI 20 ba leic SS: unkr DNA 10	ISTIC acid both lown (gen	CS: pairs i	:)	1eum	oniae	•	٠				
GAT CCT CTA TCT ATC AAT CAA CAA GGG AAT GAC CGT GGT CGC CAA TAT ASp Pro Leu Ser Ile Asn Gin Gin Gly Asn Asp Arg Gly Arg Gin Tyr 1 10 15 15 16 16 17 17 18 18 18 18 18 18 18 18 18 18 18 18 18	,		FEA (A	3) CI ATURE A) NA	ONE : : : : : :	SPF ŒY:	CDS	19										
Asp Pro Leu Ser Ile Asn Gln Gln Gly Asn Asp Arg Gly Arg Gln Tyr 15 15 1 15 1 15 1 15 1 15 1 15 1 15 1		(xi)	SEC	QUENC	E DE	SCRI	PTIC	N: 5	SEQ I	D NO	:5:							
Arg Thr Gly Ile Tyr Tyr Gln Asp Glu Ala Asp Leu Pro Ala Ile Tyr 20 25 30 Aca GTG GTG CAG GAG CAG GAA CGC ATG CTG GGT CGA AAG ATT GCA GTA Thr Val Val Gln Glu Gln Glu Arg Met Leu Gly Arg Lys Ile Ala Val 35 45	Asp				Ile					Asn					Gln			48
Thr Val Val Glu Glu Glu Glu Arg Met Leu Gly Arg Lys Ile Ala Val 35 45				Ile					Glu					Ala			•	96
GAA GTG GAG CAA TTA CGC CAC TAC ATT CTG GCT GAA GAC TAC CAC CAA 192			Val					Arg					Lys					144
	GAA	GTG	GAG	CAA	TTA	CGC	CAC	TAC	ATT	CTG	GCT	GAA	GAC	TAC	CAC	CAA		192

- 91 -

Glu	Val 50	Glu	Gln	Leu	Arg	His 55	Tyr	Ile	Leu	Ala	Glu 60	Asp	Tyr	His	Gln	
								GGT Gly								24
								GCA Ala							AGT Ser	28
								TCT Ser 105							ACA Thr	336
								TTT Phe								384
								ATT .Ile								432
								GGT Gly								480
								CAT His								520

## (2) INFORMATION FOR SEQ ID NO:6:

- (i) SEQUENCE CHARACTERISTICS:
  - (A) LENGTH: 173 amino acids
    - (B) TYPE: amino acid
    - (D) TOPOLOGY: linear
- (ii) MOLECULE TYPE: protein
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO:6:

Asp Fro Leu Ser Ile Asn Gln Gln Gly Asn Asp Arg Gly Arg Gln Tyr

10

Arg Thr Gly Ile Tyr Tyr Gln Asp Glu Ala Asp Leu Pro Ala Ile Tyr
25

Thr Val Val Gln Glu Gln Glu Arg Met Leu Gly Arg Lys Ile Ala Val
45

Glu Val Glu Gln Leu Arg His Tyr Ile Leu Ala Glu Asp Tyr His Gln
55

Asp Tyr Leu Arg Lys Asn Pro Ser Gly Tyr Cys His Ile Asp Val Thr
70

Asp Ala Asp Lys Pro Leu Ile Asp Ala Ala Asn Tyr Glu Lys Pro
95

Gln Glu Val Leu Lys Ala Ser Leu Ser Glu Glu Ser Tyr Arg Val Thr
105

105

107

Gln Glu Ala Ala Thr Glu Ala Pro Phe Thr Asn Ala Tyr Asp Gln Thr

- 92 -

		115					120					125				
Dha d			a1	710	m	Tro 1		710	The	Th-	<b>~</b> 1		Dra	T 011	Dho	
Phe (	130	GIU	GIĀ	116	ıyr	135	Asp	116	inr	Inr	140	GIU	PIO	Leu	Pne	
Phe 1 145	Ala	Lys	yab	ГÀа	Phe 150	Ala	Ser	Gly	Cys	Gly 155	Trp	Pro	Ser	Phe	Ser 160	
Arg I	Pro	Ile	Ser	Lys 165	Glu	Leu	Ile	His	Tyr 170	Tyr	Lys	Asp				
(2)	INFO	RMA	rion	FOR	SEQ	ID 1	10:7	:								
	(i)	() ()	A) LE 3) T? C) S?	CE CE ENGTE YPE: TRANI OPOLO	i: 28	32 ba leic SSS:	ase p acid	pair: d	3					•		
	(ii)	MOI	ECUI	LE TY	PE:	DNA	(ger	nomi	c)							
( ;	lii)	нун	POTHE	TIC!	AL: 1	10										
-	(iv)	ANT	ri-Si	INSE :	NO											
1	(vi)	(2	A) OF	AL SO RGANI FRAIN	:Ma	Stre	epto	coccı	ıs pı	neumo	oniae	•				
(1	rii)			ONE:												
•	(ix)	(2		3: AME/P OCATI			281									
(	(xi)	SEC	UENC	E DE	SCRI	PTIC	on: S	SEQ :	D NO	0:7:						
CC TC Se														Phe C		47
GCT G Ala A	CT	AGA Arg	GAA Glu	GGA Gly 20	CGT Arg	ACC Thr	AAT Asn	TCT Ser	GTC Val 25	CTC Leu	GGT Gly	GAA Glu	CTC Leu	GGT Gly 30	AAC Asn	95
TTC T																143
AAG G Lys A	CT	CTC Leu 50	AGC Ser	AAG Lys	GAT Asp	AAC Asn	CTC Leu 55	CTT Leu	CAG Gln	ATT Ile	GTC Val	GAG Glu 60	CTC Leu	ATG Met	CTA Leu	191
GCA G Ala A	SAT ASP 65	GTT Val	AAC Asn	AAG Lys	CGC Arg	CTC Leu 70	TCT Ser	AGT Ser	AAC Asn	AAC Asn	ATT Ile 75	CGT Arg	TTG Leu	GAT Asp	GTA Val	. 239
ACT G Thr A 80																281
С																282

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(2) INFORMATION FOR SEQ ID NO:8:

(i) SEQUENCE CHARACTERISTICS:

(A) LENGTH: 93 amino acids (B) TYPE: amino acid

(D) TOPOLOGY: linear

(ii) MOLECULE TYPE: protein

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:8:

Ser Asn Ala Gly Thr Gly Lys Thr Glu Ala Ser Val Gly Phe Gly Ala

Ala Arg Glu Gly Arg Thr Asn Ser Val Leu Gly Glu Leu Gly Asn Phe

Phe Ser Pro Glu Phe Met Asn Arg Phe Asp Gly Ile Ile Glu Phe Lys

Ala Leu Ser Lys Asp Asn Leu Leu Gln Ile Val Glu Leu Met Leu Ala

Asp Val Asn Lys Arg Leu Ser Ser Asn Asn Ile Arg Leu Asp Val Thr

Asp Lys Val Lys Glu Lys Leu Val Asp Leu Gly Tyr Asp

(2) INFORMATION FOR SEQ ID NO:9:

(i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 327 base pairs

(B) TYPE: nucleic acid

(C) STRANDEDNESS: both

(D) TOPOLOGY: unknown

(ii) MOLECULE TYPE: DNA (genomic)

(iii) HYPOTHETICAL: NO

(iv) ANTI-SENSE: NO

(vi) ORIGINAL SOURCE:

(A) ORGANISM: Streptococcus pneumoniae

(B) STRAIN: R6

(vii) IMMEDIATE SOURCE:

(B) CLONE: SPRU87

(ix) FEATURE:

(A) NAME/KEY: CDS (B) LOCATION: 3..326

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:9:

AA GTG AAA GTT GAC GAC GGC TCT CAA GCT GTA AAC ATT ATC AAC CTT Val Lys Val Asp Asp Gly Ser Gln Ala Val Asn Ile Ile Asn Lee 1  $$^{10}$ 

CTT GGT GGA CGT GTA AAC ATC GTT GAT GTT GAT GCA TGT ATG ACT CGT Leu Gly Gly Arg Val Asn Ile Val Asp Val Asp Ala Cys Met Thr Arg

47

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	CGT Arg								143
	AAA Lys								191
	GCT Ala 65								239
	ATC Ile								287
	ACT Thr						С		327

- (2) INFORMATION FOR SEO ID NO:10:
  - (i) SEQUENCE CHARACTERISTICS:
    - (A) LENGTH: 108 amino acids
    - (B) TYPE: amino acid (D) TOPOLOGY: linear
  - (ii) MOLECULE TYPE: protein
  - (xi) SEQUENCE DESCRIPTION: SEQ ID NO:10:
- Val Lys Val Asp Asp Gly Ser Gln Ala Val Asn Ile Ile Asn Leu Leu
- Gly Gly Arg Val Asn Ile Val Asp Val Asp Ala Cys Met Thr Arg Leu 25 30

15

- Arg Val Thr Val Lys Asp Ala Asp Lys Val Gly Asn Ala Glu Gln Trp
- Lys Ala Glu Gly Ala Met Gly Leu Val Met Lys Gly Gln Gly Val Gln 50 60
- Ala Ile Tyr Gly Pro Lys Ala Asp Ile Leu Lys Ser Asp Ile Gln Asp
- Ile Leu Asp Ser Gly Glu Ile Ile Pro Glu Thr Leu Pro Ser Gln Met
- Thr Glu Val Gln Gln Asn Thr Val His Phe Lys Asp 100
- (2) INFORMATION FOR SEQ ID NO:11:
  - (i) SEQUENCE CHARACTERISTICS:
    - (A) LENGTH: 417 base pairs
    - (B) TYPE: nucleic acid
    - (C) STRANDEDNESS: both
    - (D) TOPOLOGY: unknown ·
  - (ii) MOLECULE TYPE: DNA (genomic)
  - (iii) HYPOTHETICAL: NO

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(477)	ANTT-	SENSE:	MO

(vi) ORIGINAL SOURCE:

(A) ORGANISM: Streptococcus pneumoniae (B) STRAIN: R6

(vii) IMMEDIATE SOURCE: (B) CLONE: SPRU24

(ix) FEATURE:

(A) NAME/KEY: CDS (B) LOCATION: 3..416

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:11:

TT									GAC Asp		47
					Arg			Tyr	GGT		95
				Gly					ACT		143
			Leu						CCA Pro 60		191
		Ile							CTT Leu		239
	Ala								AAT Asn		287
					Phe			Ala	GTT Val		335
				Lys			Val		GCT Ala		383
			Ser				TTG Leu				417

## (2) INFORMATION FOR SEO ID NO:12:

- (i) SEQUENCE CHARACTERISTICS:
  - (A) LENGTH: 138 amino acids
    (B) TYPE: amino acid
    (D) TOPOLOGY: linear
- (ii) MOLECULE TYPE: protein
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO:12:

Ser Gln Pro Val Ser Phe Asp Thr Gly Leu Gly Asp Gly Arg Met Val

Phe Val Leu Pro Arg Glu Asn Lys Thr Tyr Phe Gly Thr Thr Asp Thr  $20 \ 25 \ 30$ 

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- 96 -10

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The Ala Ser Glu Phe Glu Leu Gly Thr Pro Leu Ser Gln Glu Lys 15  TTA GAC CAT CAC AAA CCA CAG AAA CCA TCT GAT ATT CAG GCT CTA GCC Leu Asp His His Lys Pro Gln Lys Pro Ser Asp Ile Gln Ala Leu Ala 20 25 30	Asp	Tyr	Thr 35	Gly	Asp	Leu	Glu	His 40	Pro	Lys	Val	Thr	Gln 45	Glu	Asp	Val		
Ala Gly Asn Ser Ala Ser Asp Tyr Asn Gly Gly Asn Asn Gly Thr Ile 85  Arg Asp Glu Ser Phe Asp Asn Leu Ile Ala Thr Val Glu Ser Tyr Leu 100  Ser Lys Glu Lys Thr Arg Glu Asp Val Glu Ser Ala Val Ser Lys Leu 115  Glu Ser Ser Thr Ser Glu Lys His Leu Asp 130  (2) INFORMATION FOR SEQ ID NO:13:  (i) SEQUENCE CHARACTERISTICS: (A) Length: 246 base pairs (B) TYPE: nucleic acid (C) STRANBEDWESS: both (D) TOPOLOGY: unknown  (ii) MOLECULE TYPE: DNA (genomic)  (iii) HYPOTHETICAL: NO  (iv) ANTI-SENSE: NO  (vi) ORIGINAL SOURCE: (A) ORGANISM: Streptococcus pneumonise (B) STRAIN: R6  (vii) IMMEDIATE SOURCE: (B) CLONE: SPRU75  (ix) FEATURE: (A) NAME/KEY: CDS (B) LOCATION: 3245  (xi) SEQUENCE DESCRIPTION: SEQ ID NO:13:  CG ACG GCC AGT GAA TTC GAG CTC GGT ACC CTC TC AGT CAG GAG AAA Thr Ala Ser Glu Phe Glu Leu Gly Thr Pro Leu Ser Gln Glu Lys 1  TTA GAC CAT CAC AAA CCA CAG AAA CCA TCT GAT ATT CAG GCT CTA GCC Leu Asp His His Lys Pro Gln Lys Pro Ser Asp Ile Gln Ala Leu Ala 25  TTG CTG GAA ATC TTG GAC CCC ATT CGA GAG GGG GAC GAG AGA GCTG Leu Leu Glu Ile Leu Asp Pro Sle Agg Glu Glu Ala Cal TCR Leu Leu Glu Ile Leu Asp Pro Ile Arg Glu Gly Ala Ala Glu Thr Leu  14	Asp		Leu	Leu	Gly	Ile		Asn	Asn	Arg	Phe		Glu	Ser	Asn	Ile		
Arg Asp Glu Ser Phe Asp Asn Leu Ila Ala Thr Val Glu Ser Tyr Leu 100 100 110 105 110  Ser Lys Glu Lys Thr Arg Glu Asp Val Glu Ser Ala Val Ser Lys Leu 115 120 125  Glu Ser Ser Thr Ser Glu Lys His Leu Asp 130 125  (2) INFORMATION FOR SEQ ID NO:13:  (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 246 base pairs (B) TypE: nucleic acid (C) STRANDEDENES: both (D) TOPOLOGY: unknown  (ii) MOLECULE TYPE: DNA (genomic)  (iii) HYPOTHETICAL: NO  (iv) ANTI-SENSE: NO  (vi) ORIGINAL SOURCE: (A) ORGANISM: Streptococcus pneumoniae (B) STRAIN: R6  (vii) IMMEDIATE SOURCE: (B) CLONE: SPRU75  (ix) FEATURE: (B) LOCATION: 3245  (xi) SEQUENCE DESCRIPTION: SEQ ID NO:13:  CG ACG GCC AGT GAA TYC GAG CTC GGT ACC CCT CTC AGT CAG GAG AAA Thr Ala Ser Glu Phe Glu Leu Gly Thr Pro Leu Ser Gln Glu Lys 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1		Ile	Asp	Asp	Ile		Ser	Ser	Trp	Ala		Leu	Arg	Pro	Leu			
Ser Lys Glu Lys Thr Arg Glu Asp Val Glu Ser Ala Val Ser Lys Leu 115 120 125  Glu Ser Ser Thr Ser Glu Lys His Leu Asp 130 135  (2) INFORMATION FOR SEQ ID NO:13:  (i) SEQUENCE CHARACTERISTICS: (A) Length: 246 base pairs (B) TYPE: Nucleic acid (C) STRANDEDNESS: both (D) TOPOLOGY: unknown  (ii) MOLECULE TYPE: DNA (genomic)  (iii) HYPOTHETICAL: NO  (iv) ANTI-SENSE: NO  (vi) ORIGINAL SOURCE: (A) ORGANIAM: Streptococcus pneumoniae (B) STRAIN: R6  (vii) IMMEDIATE SOURCE: (B) CLONE: SPRUT5  (ix) FEATURE: (A) NAMBEKEY: CDS (B) LOCATION: 3245  (xi) SEQUENCE DESCRIPTION: SEQ ID NO:13:  CG ACG GCC AGT GAA TTC GAG CTC GGT ACC CCT CTC AGT CAG GAG AAA Thr Ala Ser Glu Phe Glu Leu Gly Thr Pro Leu Ser Gln Glu Lys 1 1 15  TTA GAC CAT CAC AAA CCA CAG AAA CCA TCT GAT ATT CAG GCT CTA GCC Leu Asp His His Lys Pro Gln Lys Pro Ser Asp Ile Gln Ala Leu Ala TTG CTG GAA ATC TG GAC CCC ATT CGA GAG GGG GCA GAG AGA GCTG Leu Leu Glu Ile Leu Asp Pro Gln Lys Glo Gly Ala Ala Glu Thr Leu Leu Glu Ile Leu Asp Pro Tl Arg Glu Gly Ala Ala Glu Thr Leu Leu Glu Ile Leu Asp Pro Tl Arg Glu Gly Ala Ala Glu Thr Leu  14	Ala	Gly	Asn	Ser		Ser	Asp	Tyr	Asn		Gly	Asn	Asn	Gly		Ile		
Glu Ser Ser Thr Ser Glu Lys His Leu Asp 130  (2) INFORMATION FOR SEQ ID NO:13:  (i) SEQUENCE CHARACTERISTICS: (i) LENGTH: 246 base pairs (i) TYPE: nucleic acid (i) STRANDENRESS: both (ii) MOLECULE TYPE: DNA (genomic) (iii) HYPOTHETICAL: NO (iv) ANTI-SENSE: NO (vi) ORIGINAL SOURCE: (A) ORGANISM: Streptococcus pneumoniae (B) STRAIN: R6 (vii) IMMEDIATE SOURCE: (B) CLONE: SPRU75 (ix) FEATURE: (A) NAMEKEY: CDS (B) LOCATION: 3245  (xi) SEQUENCE DESCRIPTION: SEQ ID NO:13:  CG ACG GCC AGT GAA TTC GAG CTC GGT ACC CTC TCC AGT CAG GAG AAA Thr Ala Ser Glu Phe Glu Leu Gly Thr Pro Leu Ser Gln Glu Lys 1  TTA GAC CAT CAC AAA CCA CAG AAA CCA TCT GAT ATT CAG GCT CTA GCC Leu Asp His His Lys Pro Gln Lys Pro Ser Asp He Gln Ala Leu Ala 25  TTG CTG GAA ATC TTG GAC CCC ATT CGA GAG GGA GAG AGG ACC CTG Leu Leu Glu II Le Leu Asp Pro SIL Arg Glu Gly Ala Ala Glu Thr Leu  14	Arg	Asp	Glu		Phe	Asp	Asn	Leu		Ala	Thr	Val ·	Glu		Tyr	Leu		
(2) INFORMATION FOR SEQ ID NO:13:  (3) SEQUENCE CHARACTERISTICS: (A) LENGTH: 246 base pairs (B) TYPE: nucleic acid (C) STRANDEBNESS: both (D) TOPOLOGY: unknown  (ii) MOLECULE TYPE: DNA (genomic)  (iii) HYPOTHETICAL: NO  (iv) ANTI-SENSE: NO  (vi) ORIGINAL SOURCE: (A) ORGANISM: Streptococcus pneumoniae (B) STRAIN: R6  (vii) IMMEDIATE SOURCE: (B) CLONE: SPRU75  (ix) PEATURE: (A) NAME/KEY: CDS (B) LOCATION: 3245  (xi) SEQUENCE DESCRIPTION: SEQ ID NO:13:  CG ACG GCC AGT GAA TTC GAG CTC GGT ACC CCT CTC AGT CAG GAG AAA Thr Ala Ser Glu Phe Glu Leu Gly Thr Pro Leu Ser Gln Glu Lys 1  TTA GAC CAT CAC AAA CCA CAG AAA CCA TCT GAT ATT CAG GCT CTA GCC Leu Asp His His Lys Pro Gln Lys Pro Ser Asp Ile Gln Ala Leu Ala 20 25  TTG CTG GAA ATC TTG GAC CCT CTT CAG GAG GAG ACG CTG Leu Leu Glu Ile Leu Asp Pro Glu Lyg Flo Ser Asp Ile Gln Ala Leu Ala 20 25  TTG CTG GAA ATC TTG GAC CCC ATT CCA GAG GAG ACG CTG Leu Leu Glu Ile Leu Asp Pro Ile Arg Glu Gly Ala Ala Glu Thr Leu  14	Ser	Lys		Lys	Thr	Arg	Glu		Val	Glu	Ser	Ala		Ser	Lys	Leu		
(i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 246 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: both (D) TOPOLOGY: unknown  (ii) MOLECULE TYPE: DNA (genomic)  (iii) HYPOTHETICAL: NO (iv) ANTI-SENSE: NO (vi) ORIGINAL SOURCE: (A) ORGANISM: Streptococcus pneumonise (B) STRAIN: R6 (vii) IMMEDIATE SOURCE: (B) CLONE: SPRUT5 (ix) FEATURE: (A) NAMEKEY: CDS (B) LOCATION: 3245  (xi) SEQUENCE DESCRIPTION: SEQ ID NO:13:  CG ACG GCC AGT GAA TTC GAG CTC GGT ACC CCT CTC AGT CAG GAG AAA Thr Ala Ser Glu Phe Glu Leu Gly Thr Pro Leu Ser Gln Glu Lys 1  TTA GAC CAT CAC AAA CCA CAG AAA CCA TCT GAT ATT CAG GCT CTA GCC Leu Asp His His Lys Pro Gln Lys Pro Ser Asp He Gln Ala Leu Ala TTG CTG GAA ATC TTG GAC CCC ATT CGA GGG GGA GAG AGC GTG Leu Leu Glu IIe Leu Asp Pro Gln Lyg Glu Gly Ala Ala Glu Thr Leu Leu Glu IIe Leu Asp Pro Tl Agg GG GGA GGA GAG AGA CCTT Leu Leu Glu IIe Leu Asp Pro Tl Agg GL GIY Ala Ala Glu Thr Leu  14	Glu		Ser	Thr	Ser	Glu		His	Leu	Asp								
(A) LENGTH: 246 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: both (D) TOPOLOGY: unknown  (ii) MOLECULE TYPE: DNA (genomic)  (iii) HYPOTHETICAL: NO (iv) ANTI-SENSE: NO (vi) ORIGINAL SOURCE: (A) ORGANIAL SOURCE: (B) STRAIN: R6 (vii) IMMEDIATE SOURCE: (B) CLONE: SPRUTS (ix) FEATURE: (A) NAMBE/KEY: CDS (B) LOCATION: 3245  (xi) SEQUENCE DESCRIPTION: SEQ ID NO:13:  CG ACG GCC AGT GAA TTC GAG CTC GGT ACC CCT CTC AGT CAG GAG AAA Thr Ala Ser Glu Phe Glu Leu Gly Thr Pro Leu Ser Gln Glu Lys 1  TTA GAC CAT CAC AAA CCA CAG AAA CCA TCT GAT ATT CAG GCT CTA GCC Leu Asp His His Lys Pro Gln Lys Pro Ser Asp Ile Gln Ala Leu Ala 25 TTG CTG GAA ATC TTG GAC CCC ATT CGA GAG GGG GCA GAG AGC CTG TTG CTG GAA ATC TTG GAC CCC ATT CGA GGG GGA GAG ACC CTG Leu Leu Glu Ile Leu Asp Pro Sle Larg Glu Gly Ala Ala Glu Thr Leu Leu Glu Ile Leu Asp Pro Tla Arg Glu Gly Ala Ala Glu Thr Leu  14	(2)	INFO	RMAT	CION	FOR	SEQ	ID N	10:13	3:									
(iii) HYPOTHETICAL: NO  (iv) ANTI-SENSE: NO  (vi) ORIGINAL SOURCE: (A) ORGANISM: Streptococcus pneumoniae (B) STRAIN: R6  (vii) IMMEDIATE SOURCE: (B) CLONE: SPRUTS  (ix) PEATURE: (A) NAME/KEY: CDS (B) LOCATION: 3245  (xi) SEQUENCE DESCRIPTION: SEQ ID NO:13:  CG ACG GCC AGT GAA TTC GAG CTC GGT ACC CCT CTC AGT CAG GAG AAA Thr Ala Ser Glu Phe Glu Leu Gly Thr Pro Leu Ser Gln Glu Lys 1  TTA GAC CAT CAC AAA CCA CAG AAA CCT T GAT ATT CAG GCT CTA GCC Leu Asp His His Lys Pro Gln Lys Pro Ser Asp 1le Gln Ala Leu Ala 20  TTG CTG GAA ATC TTG GAC CCC ATT CGA GAG GGA GCA GAG ACG CTG Leu Leu Glu Ile Leu Asp Pro Tl GAG GAG GGA GCA GAG ACG CTG Leu Leu Glu Ile Leu Asp Pro Tl GAG GAG GGA GCA GAG ACG CTG Leu Leu Glu Ile Leu Asp Pro Tl GAG GIG GIJy Ala Ala Glu Thr Leu		(i)	(I	A) LE B) T'S C) ST	ENGTI PE: TRANI	i: 24 nucl	6 ba leic SSS:	acio both	pairs	3								
(iv) ANTI-SENSE: NO  (vi) ORIGINAL SOURCE: (A) ORGANISM: Streptococcus pneumoniae (B) STRAIN: R6  (vii) IMMEDIATE SOURCE: (B) CLONE: SPRU75  (ix) FEATURE: (A) NAME/KEY: CDS (B) LOCATION: 3245  (xi) SEQUENCE DESCRIPTION: SEQ ID NO:13:  CG ACG GCC AGT GAA TTC GAG CTC GGT ACC CCT CTC AGT CAG GAG AAA Thr Ala Ser Glu Phe Glu Leu Gly Thr Pro Leu Ser Gln Glu Lys 10  TTA GAC CAT CAC AAA CCA CAG AAA CCA TCT GAT ATT CAG GCT CTA GCC Leu Asp His His Lys Pro Gln Lys Pro Ser Asp Ile Gln Ala Leu Ala 20 25  TTG CTG GAA ATC TTG GAC CCC ATT CGA GAG GGA GCA GAG ACG CTG Leu Leu Glu Ile Leu Asp Pro Tl Arg Glu Gly Ala Ala Glu Thr Leu Leu Glu Ile Leu Asp Pro Tl Arg Glu Gly Ala Ala Glu Thr Leu		(ii)	MOI	ECUI	E T	PE:	DNA	(ger	omi	:)								
(vi) ORIGINAL SOURCE: (A) ORGANISM: Streptococcus pneumoniae (B) STRAIN: R6  (vii) IMMEDIATE SOURCE: (B) CLONE: SPRU75  (ix) FEATURE: (A) NAME/KEY: CDS (B) LOCATION: 3245  (xi) SEQUENCE DESCRIPTION: SEQ ID NO:13:  CG ACG GCC AGT GAA TTC GAG CTC GGT ACC CCT CTC AGT CAG GAG AAA Thr Ala Ser Glu Phe Glu Leu Gly Thr Pro Leu Ser Gln Glu Lys 10  TTA GAC CAT CAC AAA CCA CAG AAA CCA TTC GAT ATT CAG GCT CTA GCC Leu Asp His His Lys Pro Gln Lys Pro Ser Asp Ile Gln Ala Leu Ala 20 25  TTG CTG GAA ATC TTG GAC CCC ATT CGA GAG GAG ACG CTG Leu Leu Glu Ile Leu Asp Pro Ile Arg Glu Gly Ala Ala Glu Thr Leu Leu Glu Ile Leu Asp Pro Tle Arg Glu Gly Ala Ala Glu Thr Leu  14		(iii)	HYE	OTHE	TIC	L: N	10											
(A) ORGANISM: Streptococcus pneumoniae (B) STRAIN: R6  (vii) IMMEDIATE SOURCE: (B) CLONE: SPRU75  (ix) PEATURE: (A) NAME/KEY: CDS (B) LOCATION: 3245  (xi) SEQUENCE DESCRIPTION: SEQ ID NO:13:  CG ACG GCC AGT GAA TTC GAG CTC GGT ACC CCT CTC AGT CAG GAG AAA Thr Ala Ser Glu Phe Glu Leu Gly Thr Pro Leu Ser Gln Glu Lys 1 10  TTA GAC CAT CAC AAA CCA CAG AAA CCA TCT GAT ATT CAG GCT CTA GCC Leu Asp His His Lys Pro Gln Lys Pro Ser Asp Ile Gln Ala Leu Ala 20  TTG CTG GAA ATC TTG GAC CCT ATT CGA GAG GGA GCA GAG ACG CTG Leu Leu Glu Ile Leu Asp Pro Ile Arg Glu Gly Ala Ala Glu Thr Leu Leu Glu Ile Leu Asp Pro Tl Arg Glu Gly Ala Ala Glu Thr Leu  14		(iv)	ANT	ı-sı	NSE:	NO												
(B) CLONE: SPRU75  (ix) FEATURE: (A) NAME/KEY: CDS (B) LOCATION: 3245  (xi) SEQUENCE DESCRIPTION: SEQ ID NO:13:  CG ACG GCC AGT GAA TTC GAG CTC GGT ACC CCT CTC AGT CAG GAG AAA Thr Ala Ser Glu Phe Glu Leu Gly Thr Pro Leu Ser Gln Glu Lys 1 5 1 15  TTA GAC CAT CAC AAA CCA CAG AAA CCA TTG GAT ATT CAG GCT CTA GCC Leu Asp His His Lys Pro Gln Lys Pro Ser Asp Ile Gln Ala Leu Ala 25 30 17  TTG CTG GAA ATC TTG GAC CCC ATT CGA GAG GGA GCA GAG AGA CCTG Leu Leu Glu Ile Leu Asp Pro Ile Arg Glu Gly Ala Ala Glu Thr Leu Leu Glu Ile Leu Asp Pro Tl Arg Glu Gly Ala Ala Glu Thr Leu		(vi)	(2	) OF	(GAN	SM:	Stre	ptoc	coccu	ıs pr	eumo	niae	•					
(A) NAME/KEY: CDS (B) LOCATION: 3245  (xi) SEQUENCE DESCRIPTION: SEQ ID NO:13:  CG ACG GCC AGT GAA TTC GAG CTC GGT ACC CCT CTC AGT CAG GAG AAA Thr Ala Ser Glu Phe Glu Leu Gly Thr Pro Leu Ser Gln Glu Lys 1 10  TTA GAC CAT CAC AAA CCA CAG AAA CCA TCT GAT ATT CAG GCT CTA Leu Asp His Lys Pro Gln Lys Pro Ser Asp Ile Gln Ala Leu Ala 2 25  TTG CTG GAA ATC TTG GAC CCC ATT CGA GAG GGA GCA GAG ACG CTG Leu Leu Glu Ile Leu Asp Pro Ile Arg Glu Gly Ala Ala Glu Thr Leu  14	•	(vii)																
CG ACG GCC AGT GAA TTC GAG CTC GGT ACC CCT CTC AGT CAG GAG AAA Thr Ala Ser Glu Phe Glu Leu Gly Thr Pro Leu Ser Gln Glu Lys 1 5 10  TTA GAC CAT CAC AAA CCA CAG AAA CCA TCT GAT ATT CAG GCT CTA GCC Leu Asp His His Lys Pro Gln Lys Pro Ser Asp Ile Gln Ala Leu Ala 20  TTG CTG GAA ATC TTG GAC CCC ATT CGA GAG GGA GCA GCA GAG ACG CTG Leu Leu Glu Ile Leu Asp Pro Ile Arg Glu Gly Ala Ala Glu Thr Leu		(ix)	( A	) NA	ME/I			45										
The Ala Ser Glu Phe Glu Leu Gly Thr Pro Leu Ser Gln Glu Lys 15 15  TTA GAC CAT CAC AAA CCA CAG AAA CCA TCT GAT ATT CAG GCT CTA GCC Leu Asp His His Lys Pro Gln Lys Pro Ser Asp Ile Gln Ala Leu Ala 20 25  TTG CTG GAA ATC TTG GAC CCC ATT CGA GAG GGA GCA GCA GAG ACG CTG Leu Leu Glu Ile Leu Asp Pro Ile Arg Glu Gly Ala Ala Glu Thr Leu		(xi)	SEC	UENC	E DE	SCRI	PTIC	N: 5	EQ I	D NO	:13:							
Leu Asp His His Lys Pro Gln Lys Pro Ser Asp Ile Gln Ala Leu Ala 20 30  TTG CTG GAA ATC TTG GAC CCC ATT CGA GAG GGA GCA GCA GAG ACG CTG Leu Leu Glu Ile Leu Asp Pro Ile Arg Glu Gly Ala Ala Glu Thr Leu		hr A				he 0					ro L					ys	-	4
Leu Leu Glu Ile Leu Asp Pro Ile Arg Glu Gly Ala Ala Glu Thr Leu					Lys					Ser					Leu			9
	TTG Leu	CTG Leu	GAA Glu	Ile	TTG Leu	GAC Asp	CCC Pro	ATT Ile	Arg	GAG Glu	GGA Gly	GCA Ala	GCA Ala	Glu	ACG Thr	CTG Leu		14

PCT/US94/09942 WO 95/06732

GAC TAT CTC CGT TCT CAG GAG GTG GGA CTC AAG ATT ATC TCT GGT GAC 191 Asp Tyr Leu Arg Ser Gln Glu Val Gly Leu Lys Ile Ile Ser Gly Asp AAT CCA GTT ACG GTG TCC AGC ATT GCC CAG AAG GCT GGT TTT GCG GAC 239 Asn Pro Val Thr Val Ser Ser Ile Ala Gln Lys Ala Gly Phe Ala Asp 70 TAT CAC A 246 Tyr His (2) INFORMATION FOR SEO ID NO:14: (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 81 amino acids (B) TYPE: amino acid (D) TOPOLOGY: linear (ii) MOLECULE TYPE: protein (xi) SEQUENCE DESCRIPTION: SEQ ID NO:14: Thr Ala Ser Glu Phe Glu Leu Gly Thr Pro Leu Ser Gln Glu Lys Leu Asp His His Lys Pro Gln Lys Pro Ser Asp Ile Gln Ala Leu Ala Leu Leu Glu Ile Leu Asp Pro Ile Arg Glu Gly Ala Ala Glu Thr Leu Asp
35 40 45 Tyr Leu Arg Ser Gln Glu Val Gly Leu Lys Ile Ile Ser Gly Asp Asn Pro Val Thr Val Ser Ser Ile Ala Gln Lys Ala Gly Phe Ala Asp Tyr His (2) INFORMATION FOR SEQ ID NO:15:

- (i) SEQUENCE CHARACTERISTICS:
  - (A) LENGTH: 292 base pairs
  - (B) TYPE: nucleic acid
  - (C) STRANDEDNESS: double
  - (D) TOPOLOGY: unknown
- (ii) MOLECULE TYPE: DNA (genomic)
- (iii) HYPOTHETICAL: NO
- (iv) ANTI-SENSE: NO
- (vi) ORIGINAL SOURCE: (A) ORGANISM: Streptococcus pneumoniae
  - (B) STRAIN: R6
- (vii) IMMEDIATE SOURCE: (B) CLONE: SPRU81
- (ix) FEATURE: (A) NAME/KEY: CDS

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## (B) LOCATION: 3..290

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:15						
	1-11	CEUTIENCE	DESCRIPTION.	CEO	TD	NO.15

G						GTA Val 5										47
						Glu				Thr					ATC lle	95
					Glr				Tyr					Lys	TCC Ser	143
				Met				Ser					Gln		GCG Ala	. 191
			Phe									Leu			GGG	239
Ty											Phe				CAG Gln 95	287
	GC Cg	CA														292

## (2) INFORMATION FOR SEQ ID NO:16:

- (i) SEQUENCE CHARACTERISTICS:
  - (A) LENGTH: 96 amino acids
    (B) TYPE: amino acid
    (D) TOPOLOGY: linear
- (ii) MOLECULE TYPE: protein
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO:16:

Arg Leu Ser Trp Val Thr Pro Gly Phe Ser Gln Ser Arg Arg Cys Lys

Thr Thr Ala Ser Glu Phe Glu Leu Gly Thr Leu Arg Lys Asn Ile Gly

Leu Val Leu Gln Glu Pro Phe Leu Tyr His Gly Thr Ile Lys Ser Asn 35 40 45

Ala Phe Val Asp Ala Asp Ser Phe Ile Gln Glu Leu Pro Gln Gly Tyr 65 70 75 80

Asp Ser Pro Val Ser Glu Arg Gly Ser Ser Phe Ser Thr Gly Gln Arg

(2) INFORMATION FOR SEQ ID NO:17:

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	(i)	(1 (1	A) LI B) T	ENGT: YPE: TRAN	HARA H: 3 nuc DEDN: OGY:	42 b leic ESS:	as aci bot	pair d	s							
	(ii)	MO	LECUI	LE T	YPE:	DNA	(ge	nomi	c)							
	( <b>iii</b> )	нү	РОТН	BTIC	AL: I	NO										
	(iv	AN.	ri-si	ENSE	: NO											
	(vi)	. (2	A) OI	RGAN:	OURCI ISM: N: R	Str	epto	cocc	us pi	neum	onia	•				
	(vii)				SOUR											
	(ix)		A) NJ	AME/I	KEY:		34i							•		
	(xi)	SEC	OUBNO	CE DI	ESCR:	IPTIC	ON:	SEQ :	ID N	0:17	:					
									GCT ( Ala 1							47
									TTG Leu 25							95
									CTT Leu							143
									GAC Asp						.,	191
							Thr		CTT Leu							239
									TTT Phe					CAT His 95		287
									ACA Thr 105					TAC Tyr		335
	ATC Ile															342
(2)	TNFC	PAMSC	PTON	FOR	SEO	ID 8	iO: 1	8:								

(i) SEQUENCE CHARACTERISTICS:
(A) LENGTH: 113 amino acids
(B) TYPE: amino acid
(D) TOPOLOGY: linear

-	ı	۱	Я	u	١,

,	(11)	MOLECULE	TYPE.	protein

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:18:

Ser Ser Ile Glu Lys Gln Ile Lys Ala Leu Lys Ser Gly Ala His Ile

Val Val Gly Thr Pro Gly Arg Leu Leu Asp Leu Ile Lys Arg Lys Ala 20 25 30

Leu Lys Leu Gln Asp Ile Glu Thr Leu Ile Leu Asp Glu Ala Asp Glu 35 40 45

Met Leu Asn Met Gly Phe Leu Glu Asp Ile Glu Ala Ile Ile Ser Arg 50 60

Val Pro Glu Asn Arg Gln Thr Leu Leu Phe Ser Ala Thr Met Pro Asp 65 70 75 80

Ala Ile Lys Arg Ile Gly Val Gln Phe Met Lys Ala Pro Glu His Val 85 90 95

Arg Ile Ala Ala Lys Glu Leu Thr Thr Glu Leu Val Asp Gln Tyr Tyr 100 105 110

Ile

### (2) INFORMATION FOR SEO ID NO:19:

- (i) SEQUENCE CHARACTERISTICS:
  - (A) LENGTH: 235 base pairs
  - (B) TYPE: nucleic acid
  - (C) STRANDEDNESS: both
  - (D) TOPOLOGY: unknown
- (ii) MOLECULE TYPE: DNA (genomic)
- (iii) HYPOTHETICAL: NO
- (iv) ANTI-SENSE: NO
- (vi) ORIGINAL SOURCE:
  - (A) ORGANISM: Streptococcus pneumoniae
    (B) STRAIN: R6
    - . .
- (ix) FEATURE:
  - (A) NAME/KEY: CDS
  - (B) LOCATION: 1..234
  - (xi) SEQUENCE DESCRIPTION: SEQ ID NO:19:
- GCA TTT GTA TTT GGT CGT ACC AAA CGC CGT GTG GAT GAA TTG ACT CGT
  Ala Phe Val Phe Gly Arg Thr Lys Arg Arg Val Asp Glu Leu Thr Arg
  1 15 15
- GGT TTG AAA ATT CGT GGC TTC CGT GCA GAA GGA ATT CAT GGC GAC CTA

  96
  Gly Leu Lys Ile Arg Gly Phe Arg Ala Glu Gly Ile His Gly Asp Leu

144

GAC CAA AAC AAA CGT CTT CGT GTC CTT CGT GAC TTT AAA AAT GGC AAT

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Asp Gln Asn Lys Arg Leu Arg Val Leu Arg Asp Phe Lys Asn Gly Asn CTT GAT GTT TTG GTT GCG ACA GAC GTT GCA GCG CGT GGT TTG GAT ATT 192 Leu Asp Val Leu Val Ala Thr Asp Val Ala Ala Arg Gly Leu Asp Ile TCA GGT GTG ACC CAT GTC TAC AAC TAC GAT ATT CCA CAA GAT Ser Gly Val Thr His Val Tyr Asn Tyr Asp Ile Pro Gln Asp 234 70 235

- (2) INFORMATION FOR SEQ ID NO:20:
  - (i) SEQUENCE CHARACTERISTICS:
    - (A) LENGTH: 78 amino acids (B) TYPE: amino acid
      - (D) TOPOLOGY: linear
  - (ii) MOLECULE TYPE: protein
  - (xi) SEQUENCE DESCRIPTION: SEQ ID NO:20:

Ala Phe Val Phe Gly Arg Thr Lys Arg Arg Val Asp Glu Leu Thr Arg

Gly Leu Lys Ile Arg Gly Phe Arg Ala Glu Gly Ile His Gly Asp Leu

Asp Gln Asn Lys Arg Leu Arg Val Leu Arg Asp Phe Lys Asn Gly Asn

Leu Asp Val Leu Val Ala Thr Asp Val Ala Ala Arg Gly Leu Asp Ile

Ser Gly Val Thr His Val Tyr Asn Tyr Asp Ile Pro Gln Asp

- (2) INFORMATION FOR SEO ID NO:21:
  - (i) SEQUENCE CHARACTERISTICS:
    - (A) LENGTH: 251 base pairs
    - (B) TYPE: nucleic acid (C) STRANDEDNESS: both
  - (D) TOPOLOGY: unknown
  - (ii) MOLECULE TYPE: DNA (genomic)
  - (iii) HYPOTHETICAL: NO
  - (iv) ANTI-SENSE: NO
  - (vi) ORIGINAL SOURCE:
    - (A) ORGANISM: Streptococcus pneumoniae
    - (B) STRAIN: R6
  - (vii) IMMEDIATE SOURCE: (B) CLONE: SPRU25
  - (ix) FEATURE:
    - (A) NAME/KEY: CDS
    - (B) LOCATION: complement (2..250)

PCT/US94/09942 WO 95/06732

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:21:									
GATCTTGACT	ATGGTAAACT	ACGTAAGAAA	ATTTCCTACA	TTCCACAGAC	CATAGACTCT	60			
TTACAGGGAC	AATTATTGAT	AATCTAAAAA	TTGGTAATCC	TTCTGTTACA	TATGAGGATA	120			
TGGTGAGAGT	TTGTCGTATT	GTTGTGTATT	CATGATACGA	TTCAACGCCT	TCAAAATCGT	180			
TATGGCTCCT	TTGAGAGAGG	CGGTCAAATT	CTCGGTGGAG	AGAACACGTT	GGCTTTCGAA	240			
GCGCATCTGG	G					251			

## (2) INFORMATION FOR SEO ID NO:22:

- (i) SEQUENCE CHARACTERISTICS:
  - (A) LENGTH: 83 amino acids
    (B) TYPE: amino acid
    (D) TOPOLOGY: linear
- (ii) MOLECULE TYPE: protein
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO:22:

Pro Asp Ala Leu Arg Lys Pro Thr Cys Ser Leu His Arg Glu Phe Asp Arg Leu Ser Gln Arg Ser His Asn Asp Phe Glu Gly Val Glu Ser Tyr His Glu Tyr Thr Thr Ile Arg Gln Thr Leu Thr Ile Ser Ser Tyr Val Thr Glu Gly Leu Pro Ile Phe Arg Leu Ser Ile Ile Val Pro Val Lys

Ser Leu Trp Ser Val Glu Cys Arg Lys Phe Ser Tyr Val Val Tyr His 65 Ser Gln Asp

# (2) INFORMATION FOR SEQ ID NO:23:

- (i) SEQUENCE CHARACTERISTICS:
  - (A) LENGTH: 163 amino acids (B) TYPE: amino acid (D) TOPOLOGY: unknown
- (ii) MOLECULE TYPE: peptide
- (iii) HYPOTHETICAL: NO
- (iv) ANTI-SENSE: NO
  - (v) FRAGMENT TYPE: internal
- (vi) ORIGINAL SOURCE: (A) ORGANISM: Streptococcus pneumoniae
- (xi) SEQUENCE DESCRIPTION: SEO ID NO:23:

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Asp Arg Ser Ala Tyr Ser Ala Gln Ile Asn Gly Lys Asp Gly Ala Ala 1 10 15

Leu Ala Val Arg Asn Leu Phe Val Lys Pro Asp Phe Val Ser Ala Gly 20 25 30

Glu Lys Thr Phe Gly Asp Leu Val Ala Ala Gln Leu Pro Ala Tyr Gly 35 40 45

Asp Glu Trp Lys Gly Val Asn Leu Ala Asp Gly Gln Asp Gly Leu Phe 50 60

Asn Ala Asp Lys Ala Lys Ala Glu Phe Arg Lys Ala Lys Lys Ala Leu 65 70 75 80

Glu Ala Asp Gly Val Gln Phe Pro Ile His Leu Asp Val Pro Val Asp 85 90 95

Gln Ala Ser Lys Asn Tyr Ile Ser Arg Ile Gln Ser Phe Lys Gln Ser 100 105 110

Val Glu Thr Val Leu Gly Val Glu Asn Val Val Val Asp Ile Gln Gln 115 120 125

Met Thr Ser Asp Glu Phe Leu Asn Ile Thr Tyr Tyr Ala Ala Asn Ala 130 140

Ser Ser Glu Asp Trp Asp Val Ser Gly Gly Val Ser Trp Gly Pro Asp 145 150 155

Tyr Gln Asp

### (2) INFORMATION FOR SEO ID NO:24:

- (i) SEQUENCE CHARACTERISTICS:(A) LENGTH: 77 amino acids
  - (B) TYPE: amino acid
    (D) TOPOLOGY: unknown
- (ii) MOLECULE TYPE: peptide
- (iii) HYPOTHETICAL: NO
  (iv) ANTI-SENSE: NO
  - (v) FRAGMENT TYPE: N-terminal
- (vi) ORIGINAL SOURCE:
   (A) ORGANISM: Streptococcus pneumoniae
   (B) STRAIN: R6
- (vii) IMMEDIATE SOURCE:
   (B) CLONE: SPRU42
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO:24:

His Leu Trp Asp Ile Tyr Asn Thr Asp Glu Tyr Val Ala Tyr Pro Asp 20 25 30

Asp Glu Leu Gln Val Ala Ser Thr Ile Val Asp Val Ser Asn Gly Lys

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35

Val Ile Ala Gln Leu Gly Ala Arg His Gln Ser Ser Asn Val Ser Phe

Gly Ile Asn Gln Ala Val Glu Thr Asn Arg Asp Trp Gly 65 70 75

## (2) INFORMATION FOR SEQ ID NO:25:

- (i) SEQUENCE CHARACTERISTICS:
  - (A) LENGTH: 173 amino acids
  - (B) TYPE: amino acid
    (D) TOPOLOGY: unknown
- (ii) MOLECULE TYPE: peptide
- (iii) HYPOTHETICAL: NO
- (iv) ANTI-SENSE: NO
- (v) FRAGMENT TYPE: N-terminal
- (vi) ORIGINAL SOURCE:
  - (A) ORGANISM: Streptococcus pneumoniae
    (B) STRAIN: R6
- (vii) IMMEDIATE SOURCE:
- (B) CLONE: SPRU40
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO:25:

Asp Pro Leu Ser Ile Asn Gln Gln Gly Asn Asp Arg Gly Arg Gln Tyr

Arg Thr Gly Ile Tyr Tyr Gln Asp Glu Ala Asp Leu Pro Ala Ile Tyr

Thr Val Val Gln Glu Gln Glu Arg Met Leu Gly Arg Lys Ile Ala Val

Glu Val Glu Gln Leu Arg His Tyr Ile Leu Ala Glu Asp Tyr His Gln 50 . 55 60

Asp Tyr Leu Arg Lys Asn Pro Ser Gly Tyr Cys His Ile Asp Val Thr 65 70 75 80 80

Asp Ala Asp Lys Pro Leu Ile Asp Ala Ala Asn Tyr Glu Lys Pro Ser 85 90 95

Gln Glu Val Leu Lys Ala Ser Leu Ser Glu Glu Ser Tyr Arg Val Thr

Glu Ala Ala Thr Glu Ala Pro Phe Thr Asn Ala Tyr Asp Gln Thr 115 \$120\$

Phe Glu Glu Gly Ile Tyr Val Asp Ile Thr Thr Gly Glu Pro Leu Phe 130 135

Phe Ala Lys Asp Lys Phe Ala Ser Gly Cys Gly Trp Pro Ser Phe Ser 145 150 155 160

Arg Pro Ile Ser Lys Glu Leu Ile His Tyr Tyr Lys Asp

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## (2) INFORMATION FOR SEQ ID NO:26:

- (i) SEQUENCE CHARACTERISTICS:
  - (A) LENGTH: 175 amino acids
  - (B) TYPE: amino acid (D) TOPOLOGY: unknown
- (ii) MOLECULE TYPE: peptide
- (iii) HYPOTHETICAL: NO
- (iv) ANTI-SENSE: NO
- (v) FRAGMENT TYPE: internal
- (vi) ORIGINAL SOURCE:
  - (A) ORGANISM: Neisseria gonorrheae
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO:26:

Asp Pro Thr Ser Leu Asn Lys Gln Gly Asn Asp Thr Gly Thr Gln Tyr

Arg Ser Gly Val Tyr Tyr Thr Asp Pro Ala Glu Lys Ala Val Ile Ala

Ala Ala Leu Lys Arg Glu Gln Gln Lys Tyr Gln Leu Pro Leu Val Val 35 40 45

Glu Asn Glu Pro Leu Lys Asn Phe Tyr Asp Ala Glu Glu Tyr His Gln 50 55 60

Asp Tyr Leu Ile Lys Asn Pro Asn Gly Tyr Cys His Ile Asp Ile Arg 65 70 75 80

Lys Ala Asp Glu Pro Leu Pro Gly Lys Thr Lys Ala Ala Pro Gln Gly 85  $\phantom{\bigg|}90\phantom{\bigg|}95\phantom{\bigg|}$ 

Gln Arg Leu Arg Arg Gly Gln Arg Ile Lys Asn Arg Val Thr Pro Asn

Ser Asn Ala Pro Asp Arg Arg Ala Ile Pro Ser Asp Gln Asn Ser Ala 115 120 125

Thr Glu Tyr Ala Phe Ser His Glu Tyr Asp His Leu Phe Lys Pro Gly 130 135 140

Ile Tyr Val Asp Val Val Ser Gly Glu Pro Leu Phe Ser Ser Ala Asp 145 150 155

Lys Tyr Asp Ser Gly Cys Gly Trp Pro Ser Phe Thr Arg Pro Ile 165 170 170

- (2) INFORMATION FOR SEQ ID NO:27:
  - (i) SEQUENCE CHARACTERISTICS:(A) LENGTH: 69 amino acids
    - (B) TYPE: amino acid
    - (D) TOPOLOGY: unknown
  - (ii) MOLECULE TYPE: peptide
  - (iii) HYPOTHETICAL: NO

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- (iv) ANTI-SENSE: NO
- (v) FRAGMENT TYPE: internal
- (vi) ORIGINAL SOURCE:
  - (A) ORGANISM: Streptococcus pneumoniae
  - (B) STRAIN: R6
- (vii) IMMEDIATE SOURCE:
- (B) CLONE: SPRU39
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO:27:
- Val Leu Gly Glu Leu Gly Asn Phe Phe Ser Pro Glu Phe Met Asn Arg
- Phe Asp Gly Ile Ile Glu Phe Lys Ala Leu Ser Lys Asp Asn Leu Leu
- Gln Ile Val Glu Leu Met Leu Ala Asp Val Asn Lys Arg Leu Ser Ser
- Asn Asn Ile Arg Leu Asp Val Thr Asp Lys Val Lys Glu Lys Leu Val
- Asp Leu Gly Tyr Asp
- (2) INFORMATION FOR SEO ID NO:28:
  - (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 69 amino acids

    - (B) TYPE: amino acid
      (D) TOPOLOGY: unknown
  - (ii) MOLECULE TYPE: peptide
  - (iii) HYPOTHETICAL: NO
  - (iv) ANTI-SENSE: NO
  - (v) FRAGMENT TYPE: internal
  - (vi) ORIGINAL SOURCE:
    - (A) ORGANISM: Lycopersicon esculentum (tomato)
  - (xi) SEQUENCE DESCRIPTION: SEQ ID NO:28:
  - Val Thr Glu Glu Leu Lys Gln Tyr Phe Arg Pro Glu Phe Leu Asn Arg
  - Leu Asp Glu Met Ile Val Phe Arg Gln Leu Thr Lys Leu Glu Val Lys
  - Glu Ile Ala Asp Ile Met Leu Lys Glu Val Phe Glu Arg Leu Lys Val
  - Lys Glu Ile Glu Leu Gln Val Thr Glu Arg Phe Arg Asp Arg Val Val 50 60
  - Asp Glu Gly Tyr Asn

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### (2) INFORMATION FOR SEO ID NO:29:

- (i) SEQUENCE CHARACTERISTICS:
  - (A) LENGTH: 98 amino acids
  - (B) TYPE: amino acid (D) TOPOLOGY: unknown
- .....
- (ii) MOLECULE TYPE: peptide
- (iii) HYPOTHETICAL: NO
  - (iv) ANTI-SENSE: NO
  - (v) FRAGMENT TYPE: internal
- (vi) ORIGINAL SOURCE:
  - (A) ORGANISM: Streptococcus pneumoniae
    (B) STRAIN: R6
- (vii) IMMEDIATE SOURCE:
  - (B) CLONE: SPRU87
  - (xi) SEQUENCE DESCRIPTION: SEQ ID NO:29:
  - Asp Asp Gly Ser Gln Ala Val Asn Ile Ile Asn Leu Leu Gly Gly Arg
  - Val Asn Ile Val Asp Val Asp Ala Cys Met Thr Arg Leu Arg Val Thr 20 25 30
  - Val Lys Asp Ala Asp Lys Val Gly Asn Ala Glu Gln Trp Lys Ala Glu 35  $\phantom{-}40\phantom{0}$
  - Gly Ala Met Gly Leu Val Met Lys Gly Gln Gly Val Gln Ala Ile Tyr 50 60
  - Gly Pro Lys Ala Asp Ile Leu Lys Ser Asp Ile Gln Asp Ile Leu Asp 65 70 75 80
  - Ser Gly Glu Ile Ile Pro Glu Thr Leu Pro Ser Gln Met Thr Glu Val 85 90 95

Gln Gln

## (2) INFORMATION FOR SEQ ID NO:30:

- (i) SEQUENCE CHARACTERISTICS:
  - (A) LENGTH: 97 amino acids
  - (B) TYPE: amino acid (D) TOPOLOGY: unknown
- (ii) MOLECULE TYPE: peptide
- (iii) HYPOTHETICAL: NO
- (iv) ANTI-SENSE: NO
- (v) FRAGMENT TYPE: internal
- (vi) ORIGINAL SOURCE:
   (A) ORGANISM: Bacillus subtilis

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(xi) SEQUENCE DESCRIPTION: SEQ ID NO:30:

Glu Ala Gly Asp Leu Pro Tyr Glu Ile Leu Gln Ala Met Gly Asp Gln
1 5 10 15

Glu Asn Ile Lys His Leu Asp Ala Cys Ile Thr Arg Leu Arg Val Thr  $20 \\ 25 \\ 30$ 

Val Asn Asp Gln Lys Lys Val Asp Lys Asp Arg Leu Lys Gln Leu Gly

Ala Ser Gly Val Leu Glu Val Gly Asn Asn Ile Gln Ala Ile Phe Gly 50 55 60

Pro Arg Ser Asp Gly Leu Lys Thr Gln Met Gln Asp Ile Ile Ala Gly 65  $\phantom{\bigg|}70\phantom{\bigg|}$  70  $\phantom{\bigg|}80\phantom{\bigg|}$ 

Arg Lys Pro Arg Pro Glu Pro Lys Thr Ser Ala Gln Glu Glu Val Gly 85 90 . 95

Gln

#### (2) INFORMATION FOR SEQ ID NO:31:

- (i) SEQUENCE CHARACTERISTICS:
  - (A) LENGTH: 69 amino acids
  - (B) TYPE: amino acid
    (D) TOPOLOGY: unknown
- .
- (ii) MOLECULE TYPE: peptide
- (iii) HYPOTHETICAL: NO
- (iv) ANTI-SENSE: NO
- (v) FRAGMENT TYPE: internal
- (vi) ORIGINAL SOURCE:
  - (A) ORGANISM: Streptococcus pneumoniae (B) STRAIN: R6
- (vii) IMMEDIATE SOURCE:
- (B) CLONE: SPRU24
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO:31:

Asp Gly Arg Met Val Phe Val Leu Pro Arg Glu Asn Lys Thr Tyr Phe 1  $\phantom{\bigg|}$  5  $\phantom{\bigg|}$  10  $\phantom{\bigg|}$  .

Thr Gln Glu Asp Val Asp Tyr Leu Leu Gly Ile Val Asn Asn Arg Phe 35 40 45

Pro Glu Ser Asn Ile Thr Ile Asp Asp Ile Glu Ser Ser Trp Ala Gly 50 60

Leu Arg Pro Leu Ile 65

(2) INFORMATION FOR SEQ ID NO:32:

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- (i) SEQUENCE CHARACTERISTICS:
  - (A) LENGTH: 69 amino acids (B) TYPE: amino acid
    - (D) TOPOLOGY: unknown
- (ii) MOLECULE TYPE: peptide
- (iii) HYPOTHETICAL: NO
- (iv) ANTI-SENSE: NO
- (v) FRAGMENT TYPE: internal
- (vi) ORIGINAL SOURCE: (A) ORGANISM: Bacillus subtilis
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO:32:

Asp Gly Arg Met Val Phe Ala Ile Pro Arg Glu Gly Lys Thr Tyr Val

Gly Thr Thr Asp Thr Val Tyr Lys Glu Ala Leu Glu His Pro Arg Met

Thr Thr Glu Asp Arg Asp Tyr Val Ile Lys Ser Ile Asn Tyr Met Phe

Pro Glu Leu Asn Ile Thr Ala Asn Asp Ile Glu Ser Ser Trp Ala Gly

Leu Arg Pro Leu Ile

- (2) INFORMATION FOR SEQ ID NO:33:
  - (i) SEQUENCE CHARACTERISTICS:
    - (A) LENGTH: 41 amino acids (B) TYPE: amino acid
    - (D) TOPOLOGY: unknown
  - (ii) MOLECULE TYPE: peptide
  - (iii) HYPOTHETICAL: NO
  - (iv) ANTI-SENSE: NO
  - (v) FRAGMENT TYPE: internal
  - (vi) ORIGINAL SOURCE:
    - (A) ORGANISM: Streptococcus pneumoniae
    - (B) STRAIN: R6
  - (vii) IMMEDIATE SOURCE: (B) CLONE: SPRU75
  - (xi) SEQUENCE DESCRIPTION: SEQ ID NO:33:

Ala Leu Leu Glu Ile Leu Asp Pro Val Arg Glu Gly Ala Ala Glu Thr

Leu Asp Tyr Leu Arg Ser Gln Glu Val Gly Leu Lys Ile Ile Ser Gly

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Val Asn Pro Val Thr Val Ser Ser Ile 35 40

- (2) INFORMATION FOR SEQ ID NO:34:
  - (i) SEQUENCE CHARACTERISTICS:
    - (A) LENGTH: 41 amino acids
      - (B) TYPE: amino acid
        (D) TOPOLOGY: unknown
  - (ii) MOLECULE TYPE: peptide
  - (iii) HYPOTHETICAL: NO
  - (iv) ANTI-SENSE: NO
  - (v) FRAGMENT TYPE: internal
  - (vi) ORIGINAL SOURCE:
    - (A) ORGANISM: Streptococcus typhimurium
  - (xi) SEQUENCE DESCRIPTION: SEQ ID NO:34:
  - Gly Met Leu Thr Phe Leu Asp Pro Pro Lys Glu Ser Ala Gly Lys Ala
  - Ile Ala Ala Leu Arg Asp Asn Gly Val Ala Val Lys Val Leu Thr Gly 20 25 30

Asp Asn Pro Val Val Thr Ala Arg Ile 35 40

- (2) INFORMATION FOR SEQ ID NO:35:
  - (i) SEQUENCE CHARACTERISTICS:
    - (A) LENGTH: 72 amino acids
    - (B) TYPE: amino acid (D) TOPOLOGY: unknown
  - (ii) MOLECULE TYPE: peptide
  - (iii) HYPOTHETICAL: NO
  - (iv) ANTI-SENSE: NO
  - (v) FRAGMENT TYPE: internal
  - (vi) ORIGINAL SOURCE:
    - (A) ORGANISM: Streptococcus pneumoniae
      - (B) STRAIN: R6
  - (vii) IMMEDIATE SOURCE:
    - (B) CLONE: SPRU81
  - (xi) SEQUENCE DESCRIPTION: SEQ ID NO:35:
  - Gly Thr Leu Arg Lys Asn Ile Gly Leu Val Leu Gln Glu Pro Phe Leu

    1 10 15
  - Tyr His Gly Thr Ile Lys Ser Asn Ile Ala Met Tyr Gln Glu Ile Ser
  - Asp Glu Gln Val Gln Ala Ala Ala Ala Phe Val Asp Ala Asp Ser Phe

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5 40

Ile Gln Glu Leu Pro Gln Gly Tyr Asp Ser Pro Val Ser Glu Arg Gly

Ser Ser Phe Ser Thr Gly Gln Arg

- (2) INFORMATION FOR SEQ ID NO:36:
  - (i) SEQUENCE CHARACTERISTICS:
    - (A) LENGTH: 73 amino acids
      (B) TYPE: amino acid
    - (D) TOPOLOGY: unknown
  - (ii) MOLECULE TYPE: peptide
  - (iii) HYPOTHETICAL: NO
  - (iv) ANTI-SENSE: NO
  - (v) FRAGMENT TYPE: internal
  - (vi) ORIGINAL SOURCE:
    - (A) ORGANISM: Bordetella pertussis
  - (xi) SEQUENCE DESCRIPTION: SEQ ID NO:36:

Ala Ser Leu Arg Arg Gln Leu Gly Val Val Leu Gln Glu Ser Thr Leu

Phe Asn Arg Ser Val Arg Asp Asn Ile Ala Leu Thr Arg Pro Gly Ala  $20 \hspace{1.5cm} 25 \hspace{1.5cm} 30$ 

Ser Met His Glu Val Val Ala Ala Ala Arg Leu Ala Gly Ala His Glu 35 40 45

Phe Ile Cys Gln Leu Pro Glu Gly Tyr Asp Thr Met Leu Gly Glu Asn 50 60

Gly Val Gly Leu Ser Gly Gly Gln Arg

- (2) INFORMATION FOR SEO ID NO:37:
  - (i) SEQUENCE CHARACTERISTICS:
    - (A) LENGTH: 86 amino acids
    - (B) TYPE: amino acid
      (D) TOPOLOGY: unknown
  - (ii) MOLECULE TYPE: peptide
  - (iii) HYPOTHETICAL: NO
  - (iv) ANTI-SENSE: NO
  - (v) FRAGMENT TYPE: internal
  - (vi) ORIGINAL SOURCE:
    - (A) ORGANISM: Streptococcus pneumoniae (B) STRAIN: R6
  - (vii) IMMEDIATE SOURCE: (B) CLONE: SPRU17

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- (xi) SEQUENCE DESCRIPTION: SEQ ID NO:37:
- Gln Ile Lys Ala Leu Lys Ser Gly Ala His Ile Val Val Gly Thr Pro 1 5 10 15
- Gly Arg Leu Leu Asp Leu Ile Lys Arg Lys Ala Leu Lys Leu Gln Asp 20 25 30
- Ile Glu Thr Leu Ile Leu Asp Glu Ala Asp Glu Met Leu Asn Met Gly 35 40 45
- Phe Leu Glu Asp Ile Glu Ala Ile Ile Ser Arg Val Pro Glu Asn Arg 50 55 60
- Gln Thr Leu Leu Phe Ser Ala Thr Met Pro Asp Ala Ile Lys Arg Ile 65 70 75 80
- Gly Val Gln Phe Met Lys 85
- (2) INFORMATION FOR SEQ ID NO:38:
  - (i) SEQUENCE CHARACTERISTICS:
    - (A) LENGTH: 86 amino acids
    - (B) TYPE: amino acid
      (D) TOPOLOGY: unknown
  - (ii) MOLECULE TYPE: peptide
  - (iii) HYPOTHETICAL: NO
  - (iv) ANTI-SENSE: NO
  - (v) FRAGMENT TYPE: internal
  - (vi) ORIGINAL SOURCE:
    - (A) ORGANISM: Escherichia coli
  - (xi) SEQUENCE DESCRIPTION: SEQ ID NO:38:
  - Gln Leu Arg Ala Leu Arg Gln Gly Pro Gln Ile Val Val Gly Thr Pro
  - Gly Arg Leu Leu Asp His Leu Lys Arg Gly Thr Leu Asp Leu Ser Lys 20 25 30
  - Leu Ser Gly Leu Val Leu Asp Glu Ala Asp Glu Met Leu Arg Met Gly 35 40 45
  - Phe Ile Glu Asp Val Glu Thr Ile Met Ala Gln Ile Pro Glu Gly His 50 60
  - Gln Thr Ala Leu Phe Ser Ala Thr Met Pro Glu Ala Ile Arg Arg Ile 65 70 75 80
  - Thr Arg Arg Phe Met Lys
- (2) INFORMATION FOR SEQ ID NO:39:
  - (i) SEQUENCE CHARACTERISTICS:

    (A) LENGTH: 78 amino acids
    (B) TYPE: amino acid

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- (D) TOPOLOGY: unknown
- (ii) MOLECULE TYPE: peptide
- (iii) HYPOTHETICAL: NO
- (iv) ANTI-SENSE: NO
- (v) FRAGMENT TYPE: internal
- (vi) ORIGINAL SOURCE:
  - (A) ORGANISM: Escherichia coli
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO:39:

Ala Ile Ile Phe Val Arg Thr Lys Asn Ala Thr Leu Glu Val Ala Glu

Ala Leu Glu Arg Asn Gly Tyr Asn Ser Ala Ala Leu Asn Gly Asp Met 20 25 30

Asn Gln Ala Leu Arg Glu Gln Thr Leu Glu Arg Leu Lys Asp Gly Arg 35 40 40

Leu Asp Ile Leu Ile Ala Thr Asp Val Ala Ala Arg Gly Leu Asp Val 50 55 60

30

Glu Arg Ile Ser Leu Val Val Asn Tyr Asp Ile Pro Met Asp 65 70 75

- (2) INFORMATION FOR SEQ ID NO:40:
  - (i) SEQUENCE CHARACTERISTICS:
    - (A) LENGTH: 30 base pairs
    - (B) TYPE: nucleic acid (C) STRANDEDNESS: single
    - (D) TOPOLOGY: unknown
  - (ii) MOLECULE TYPE: DNA (genomic)
  - (iii) HYPOTHETICAL: NO
  - (iv) ANTI-SENSE: NO
  - (vi) ORIGINAL SOURCE:
  - (A) ORGANISM: Streptococcus pneumoniae
  - (xi) SEQUENCE DESCRIPTION: SEQ ID NO:40:

## AAAGGATCCA TGAARAARAA YMGHGTNTTY

(2) INFORMATION FOR SEQ ID NO:41:

- (i) SEQUENCE CHARACTERISTICS:
  - (A) LENGTH: 30 base pairs (B) TYPE: nucleic acid
    - (C) STRANDEDNESS: single
  - (D) TOPOLOGY: unknown
- (ii) MOLECULE TYPE: DNA (genomic)
- (iii) HYPOTHETICAL: NO

WO 95/06732		PCT/US94/09942
	- 114 -	
(iv)	ANTI-SENSE: NO	•
(vi)	ORIGINAL SOURCE: (A) ORGANISM: Streptococcus pneumoniae	
(xi)	SEQUENCE DESCRIPTION: SEQ ID NO:41:	
TTTGGATC	CG TTGGTTTAGC AAAATCGCTT	30
(2) INFO	RMATION FOR SEQ ID NO:42:	
(i)	SEQUENCE CHARACTERISTICS: (A) LENGTH: 15 base pairs (B) TYPE: nucleic acid (C) STRANDENBSS: single (D) TOPOLOGY: unknown	
· (ii)	MOLECULE TYPE: DNA (genomic)	
(iii)	HYPOTHETICAL: NO	
(iv)	ANTI-SENSE: NO	
(vi)	ORIGINAL SOURCE: (A) ORGANISM: Streptococcus pneumoniae	
(xi)	SEQUENCE DESCRIPTION: SEQ ID NO:42:	
AATATCGC	CC TGAGC	15
(2) INFO	RMATION FOR SEQ ID NO:43:	
(i)	SEQUENCE CHARACTERISTICS: (A) LENGTH: 17 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: unknown	
(ii)	MOLECULE TYPE: DNA (genomic)	
(iii)	HYPOTHETICAL: NO	
(iv)	ANTI-SENSE: NO	
(vi)	ORIGINAL SOURCE: (A) ORGANISM: Streptococcus pneumoniae	
(xi)	SEQUENCE DESCRIPTION: SEQ ID NO:43:	
ATCACGCA	GA GCGGCAG	. 17
(2) INFO	RMATION FOR SEQ ID NO:44:	

(i) SEQUENCE CHARACTERISTICS:
(A) LENGTH: 52 amino acids
(B) TYPE: amino acid
(D) TOPCOLOGY: unknown
(ii) MOLECULE TYPE: peptide
(iii) HYPOTHETICAL: NO

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- (iv) ANTI-SENSE: NO
- (v) FRAGMENT TYPE: N-terminal
- (vi) ORIGINAL SOURCE: (A) ORGANISM: Streptococcus pneumoniae
- (xi) SEQUENCE DESCRIPTION: SEO ID NO:44:

Met Lys His Leu Leu Ser Tyr Phe Lys Pro Tyr Ile Lys Glu Ser Ile

Leu Ala Pro Leu Phe Lys Leu Leu Glu Ala Val Phe Glu Leu Leu Val

Pro Met Val Ile Ala Gly Ile Val Asp Gln Ser Leu Pro Gln Gly Asp

Pro Arg Val Pro

- (2) INFORMATION FOR SEO ID NO:45:
  - (i) SEQUENCE CHARACTERISTICS:
    - (A) LENGTH: 56 amino acids (B) TYPE: amino acid
    - (D) TOPOLOGY: unknown
  - (ii) MOLECULE TYPE: peptide
  - (iii) HYPOTHETICAL: NO
  - (iv) ANTI-SENSE: NO
  - (v) FRAGMENT TYPE: N-terminal
  - (vi) ORIGINAL SOURCE:
    - (A) ORGANISM: Streptococcus pneumoniae
  - (xi) SECUENCE DESCRIPTION: SEC ID NO:45:

Met Ala Lys Asn Asn Lys Val Ala Val Val Thr Thr Val Pro Ser Val

Ala Glu Gly Leu Lys Asn Val Asn Gly Val Asn Phe Asp Tyr Lys Asp

Glu Ala Ser Ala Lys Glu Ala Ile Lys Glu Glu Lys Leu Lys Gly Tyr

Leu Thr Ile Asp Pro Arg Val Pro

- (2) INFORMATION FOR SEQ ID NO:46:
  - (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 2019 base pairs
    (B) TYPE: nucleic acid
    (C) STRANDEDNESS: both

    - (D) TOPOLOGY: linear
  - (ii) MOLECULE TYPE: CDNA

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- (iii) HYPOTHETICAL: NO
- (iv) ANTI-SENSE: NO
- (vi) ORIGINAL SOURCE:
   (A) ORGANISM: Streptococcus pneumoniae
   (B) STRAIN: R6
- (vii) IMMEDIATE SOURCE: (B) CLONE: SPRU98
- (ix) FEATURE: (A) NAME/KEY: CDS (B) LOCATION: 1..1932
- ·

(xi	) SE	QUEN	CE D	ESCR	IPTI	ON:	SEQ	ID N	0:46	:			
GTA Val												GAG Glu	48
ACA Thr													96
ACA Thr													144
GGT Gly 50													192
GAG Glu													240
CGT Arg													288
GTC Val													336
AAA Lys													384
GCC Ala 130													432
GCT Ala													480
TTC Phe													528
GAG Glu													576

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	AGT Ser															624
	AAA Lys 210															672
	AAT Asn															720
	ACC Thr															768
	CGT Arg						Ala									816
	AAG Lys															864
	GGT Gly 290															912
	GAT Asp															960
	CGT Arg															1008
	TTG Leu											Arg				1056
	CCA Pro															1104
	AAA Lys 370															1152
	GCA Ala															1200
GAA Glu	TTT Phe	GCT Ala	AAA Lys	GCT Ala 405	AAA Lys	TCA Ser	GCC Ala	TTA Leu	CAA Gln 410	GCA Ala	GAA Glu	GGT Gly	GTG Val	ACA Thr 415	TTC Phe	1248
CCA Pro	ATT Ile	CAT His	TTG Leu 420	GAT Asp	ATG Met	CCA Pro	GTT Val	GAC Asp 425	CAG Gln	ACA Thr	GCA Ala	ACT Thr	ACA Thr 430	AAA Lys	GTT Val	1296
	CGC Arg															1344
GAT	AAT	GTC	ATT	ATT	GAT	ATT	CAA	CAA	CTA	CAA	AAA	GAC	GAA	GTA	AAC	1392

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Asp	Asn 450	Val	Ile	Ile	Asp	Ile 455	Gln	Gln	Leu	Gln	Lys 460	Asp	Glu	Val	Asn	
					GCT Ala 470											1440
TCA Ser	GAT Asp	AAT Asn	GTC Val	GGT Gly 485	TGG Trp	GGT Gly	CCA Pro	GAC Asp	TTT Phe 490	GCC Ala	GAT Asp	CCA Pro	TCA Ser	ACC Thr 495	TAC Tyr	1488
CTT Leu	GAT Asp	ATC Ile	ATC Ile 500	AAA Lys	CCA Pro	TCT Ser	GTA Val	GGA Gly 505	GAA Glu	AGT Ser	ACT Thr	AAA Lys	ACA Thr 510	TAT Tyr	TTA Leu	1536
GGG Gly	TTT Phe	GAC Asp 515	TCA Ser	GGG Gly	GAA Glu	gat Asp	AAT Asn 520	GTA Val	GCT Ala	GCT Ala	AAA Lys	AAA Lys 525	GTA Val	GGT Gly	CTA Leu	1584
					TTG Leu											1632
GTT Val 545	GCT Ala	AAA Lys	CGC Arg	TAT Tyr	GAT Asp 550	AAA Lys	TAC Tyr	GCT Ala	GCA Ala	GCC Ala 555	CAA Gln	GCT Ala	TGG Trp	TTG Leu	ACA Thr 560	1680
			Leu		ATT Ile											1728
					CCA Pro											1776
					CCA Pro											1824
					GAT Asp											1872
					TCT Ser 630											1920
	GTG Val		TAAC	TGT	rgc a	TAAAA	ATA	AG AZ	AGG2	TTT	GT2	ATTTO	тст			1969
TGA	TGCT	GA I	TCCT	TIT	T AC	ATT	GTA	A AGA	AAGZ	TTC	TAAI	TGT	CT			2019

## (2) INFORMATION FOR SEQ ID NO:47:

(i) SEQUENCE CHARACTERISTICS:
(A) LERGTH: 643 amino acids
(B) TYPE: amino acid
(D) TOPOLOGY: linear

(ii) MOLECULE TYPE: protein

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:47:

Gly Val Leu Ala Ala Cys Ser Gly Ser Gly Ser Ser Ala Lys Gly Glu Lys Thr Phe Ser Tyr Ile Tyr Glu Thr Asp Pro Asp Asn Leu Asn Tyr 20 25 30 Leu Thr Thr Ala Lys Ala Ala Thr Ala Asn Ile Thr Ser Asn Val Val Asp Gly Leu Leu Glu Asn Asp Arg Tyr Gly Asn Phe Val Pro Ser Met 50 55 60 Ala Glu Asp Trp Ser Val Ser Lys Asp Gly Leu Thr Tyr Thr Tyr Thr 65 70 75 80 Ile Arg Lys Asp Ala Lys Trp Tyr Thr Ser Glu Gly Glu Glu Tyr Ala 85 90 95 Ala Val Lys Ala Gln Asp Phe Val Thr Gly Leu Lys Tyr Ala Ala Asp Lys Lys Ser Asp Ala Leu Tyr Pro Val Gln Glu Ser Ile Lys Gly Leu 115 120 125 Asp Ala Tyr Val Lys Gly Glu Ile Lys Asp Phe Ser Gln Val Gly Ile 130 135 140 Lys Ala Leu Asp Glu Gln Thr Val Gln Tyr Thr Leu Asn Lys Pro Glu 145 150 160 Ser Phe Trp Asn Ser Lys Thr Thr Met Gly Val Leu Ala Pro Val Asn 165 170 175 Glu Glu Phe Leu Asn Ser Lys Gly Asp Asp Phe Ala Lys Ala Thr Asp 180 185 190 Pro Ser Ser Leu Leu Tyr Asn Gly Pro Tyr Leu Leu Lys Ser Ile Val Thr Lys Ser Ser Val Glu Phe Ala Lys Asn Pro Asn Tyr Trp Asp Lys 210 215 220 Asp Asn Val His Ile Asp Lys Val Lys Leu Ser Phe Trp Asp Gly Gln 225 230 235 240 Asp Thr Ser Lys Pro Ala Glu Asn Phe Lys Asp Gly Ser Leu Thr Ala 245 250 255 Ala Arg Leu Tyr Pro Thr Ser Ala Ser Phe Ala Glu Leu Glu Lys Ser 260 265 270Met Lys Asp Asn Ile Val Tyr Thr Gln Gln Asp Ser Ile Thr Tyr Leu 275 280 285 Val Gly Thr Asn Ile Asp Arg Gln Ser Tyr Lys Tyr Thr Ser Lys Thr 290 295 300 Ser Asp Glu Gln Lys Ala Ser Thr Lys Lys Ala Leu Leu Asn Lys Asp 305 310 315 Phe Arg Gln Ala Ile Ala Phe Gly Phe Asp Arg Thr Ala Tyr Ala Ser 325 330 335Gln Leu Asn Gly Gln Thr Gly Ala Ser Lys Ile Leu Arg Asn Leu Phe 340 345 350

Val Pro Pro Thr Phe Val Gln Ala Asp Gly Lys Asn Phe Gly Asp Met Val Lys Glu Lys Leu Val Thr Tyr Gly Asp Glu Trp Lys Asp Val Asn 370 375 380 Leu Ala Asp Ser Gln Asp Gly Leu Tyr Asn Pro Glu Lys Ala Lys Ala 385 Glu Phe Ala Lys Ala Lys Ser Ala Leu Gln Ala Glu Gly Val Thr Phe
405 410 415 Pro Ile His Leu Asp Met Pro Val Asp Gln Thr Ala Thr Thr Lys Val Gln Arg Val Gln Ser Met Lys Gln Ser Leu Glu Ala Thr Leu Gly Ala Asp Asn Val Ile Ile Asp Ile Gln Gln Leu Gln Lys Asp Glu Val Asn Asn Ile Thr Tyr Phe Ala Glu Asn Ala Ala Gly Glu Asp Trp Asp Leu Ser Asp Asn Val Gly Trp Gly Pro Asp Phe Ala Asp Pro Ser Thr Tyr Leu Asp Ile Ile Lys Pro Ser Val Gly Glu Ser Thr Lys Thr Tyr Leu 500 505 510 Gly Phe Asp Ser Gly Glu Asp Asn Val Ala Ala Lys Lys Val Gly Leu 515 520 525 Tyr Asp Tyr Glu Lys Leu Val Thr Glu Ala Gly Asp Glu Thr Thr Asp 530 535 540Val Ala Lys Arg Tyr Asp Lys Tyr Ala Ala Ala Gln Ala Trp Leu Thr 545 550 555 560 Asp Ser Ala Leu Ile Ile Pro Thr Thr Ser Arg Thr Gly Arg Pro Ile Leu Ser Lys Met Val Pro Phe Thr Ile Pro Phe Ala Leu Ser Gly Asn 585 Lys Gly Thr Ser Glu Pro Val Leu Tyr Lys Tyr Leu Glu Leu Gln Asp Lys Ala Val Thr Val Asp Glu Tyr Gln Lys Ala Gln Glu Lys Trp Met Lys Glu Lys Glu Glu Ser Asn Lys Lys Ala Gln Glu Asp Leu Ala Lys 625 630 635 640 His Val Lys

- (2) INFORMATION FOR SEO ID NO:48:
  - (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 642 amino acids
    - (B) TYPE: amino acid
    - (D) TOPOLOGY: unknown

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(ii) MOLECULE TYPE: peptide
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- (iii) HYPOTHETICAL: NO
- .
- (iv) ANTI-SENSE: NO
- (vi) ORIGINAL SOURCE: (A) ORGANISM: Streptococcus pneumoniae
- (vii) IMMEDIATE SOURCE:
  - (B) CLONE: amiA
  - (x) PUBLICATION INFORMATION:
    - (A) AUTHORS: Alloing, et al.
      - (C) JOURNAL: Mol. Microbiol.
      - (D) VOLUME: 4
      - (F) PAGES: 633-644
    - (G) DATE: 1990
- note: the reference contains a sequence error; the correct sequence shown below is obtained from  ${\tt GENBANK}$ 
  - (xi) SEQUENCE DESCRIPTION: SEQ ID NO:48:
  - Gly Val Leu Ala Ala Cys Ser Ser Ser Lys Ser Ser Asp Ser Ser Ala 1 10 15
  - Pro Lys Ala Tyr Gly Tyr Val Tyr Thr Ala Asp Pro Glu Thr Leu Asp 20 25 30
  - Tyr Leu Ile Ser Arg Lys Asn Ser Thr Thr Val Val Thr Ser Asn Gly  ${\bf 35}$  40 45
  - Ile Asp Gly Leu Phe Thr Asn Asp Asn Tyr Gly Asn Leu Ala Pro Ala 50 60
  - Val Ala Glu Asp Trp Glu Val Ser Lys Asp Gly Leu Thr Tyr Thr Tyr 65  $\phantom{-}70\phantom{0}$  75  $\phantom{-}80\phantom{0}$
  - Lys Ile Arg Lys Gly Val Lys Trp Phe Thr Ser Asp Gly Glu Glu Tyr 85 90 95

  - Asp Lys Lys Ser Glu Ala Met Tyr Leu Ala Glu Asn Ser Val Lys Gly 115 120 125
  - Leu Ala Asp Tyr Leu Ser Gly Thr Ser Thr Asp Phe Ser Thr Val Gly 130 140
  - Val Lys Ala Val Asp Asp Tyr Thr Leu Gln Tyr Thr Leu Asn Gln Pro 145 155 160
  - Glu Pro Phe Trp Asn Ser Lys Leu Thr Tyr Ser Ile Phe Trp Pro Leu 165 170 175
  - Asn Glu Glu Phe Glu Thr Ser Lys Gly Ser Asp Phe Ala Lys Pro Thr 180 185 190
  - Asp Pro Thr Ser Leu Leu Tyr Asn Gly Pro Phe Leu Leu Lys Gly Leu 195 200 205
  - Thr Ala Lys Ser Ser Val Glu Phe Val Lys Asn Glu Gln Tyr Trp Asp 210 215

Lys Glu Asn Val His Leu Asp Thr Ile Asn Leu Ala Tyr Tyr Asp Gly 225 230 235 240 Ser Asp Gln Glu Ser Leu Glu Arg Asn Phe Thr Ser Gly Ala Tyr Ser Tyr Ala Arg Leu Tyr Pro Thr Ser Ser Asn Tyr Ser Lys Val Ala Glu 260 265 270 Glu Tyr Lys Asp Asn Ile Tyr Tyr Thr Gln Ser Gly Ser Gly Ile Ala 275 280 285 Gly Leu Gly Val Asn Ile Asp Arg Gln Ser Tyr Asn Tyr Thr Ser Lys 290 295 300 Thr Thr Asp Ser Glu Lys Val Ala Thr Lys Lys Ala Leu Leu Asn Lys 305 310 315 320 Asp Phe Arg Gln Ala Leu Asn Phe Ala Leu Asp Arg Ser Ala Tyr Ser 325 330 335 Ala Gln Ile Asn Gly Lys Asp Gly Ala Ala Leu Ala Val Arg Asn Leu 340 350 Phe Val Lys Pro Asp Phe Val Ser Ala Gly Glu Lys Thr Phe Gly Asp 355 360 365 Leu Val Ala Ala Gln Leu Pro Ala Tyr Gly Asp Glu Trp Lys Gly Val 370 380 Asn Leu Ala Asp Gly Gln Asp Gly Leu Phe Asn Ala Asp Lys Ala Lys 385 390 395 Ala Glu Phe Arg Lys Ala Lys Lys Ala Leu Glu Ala Asp Gly Val Gln 405 410 415 Phe Pro Ile His Leu Asp Val Pro Val Asp Gln Ala Ser Lys Asn Tyr 420 425 430 Ile Ser Arg Ile Gln Ser Phe Lys Gln Ser Val Glu Thr Val Leu Gly
435 440 445 Val Glu Asn Val Val Val Asp Ile Gln Gln Met Thr Ser Asp Glu Phe 450 455 460 Leu Asn Ile Thr Tyr Tyr Ala Ala Asn Ala Ser Ser Glu Asp Trp Asp 465 470 475 480 Val Ser Gly Gly Val Ser Trp Gly Pro Asp Tyr Gln Asp Pro Ser Thr 485 490 495 Tyr Leu Asp Ile Leu Lys Thr Thr Ser Ser Glu Thr Thr Lys Thr Tyr 500 505 510Leu Gly Phe Asp Asn Pro Asn Ser Pro Ser Val Val Gln Val Gly Leu 515 520 525 Lys Glu Tyr Asp Lys Leu Val Asp Glu Ala Ala Lys Glu Thr Ser Asp 530 540 Phe Asn Val Arg Tyr Glu Lys Tyr Ala Ala Ala Gln Ala Trp Leu Thr 545 550 555 560 Asp Ser Ser Leu Phe Ile Pro Ala Met Ala Ser Ser Gly Ala Ala Pro 565 570 575

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Val	Leu	Ser	Arg 580	Ile	Val	Pro	Phe	Thr 585	Gly	Ala	Ser	Ala	Gln 590	Thr	Gly
Ser	Lys	Gly 595	Ser	, qe	Val	Tyr	Phe 600	Ļys	Tyr	Leu	Lys	Leu 605	Gln	Asp	Lys
Ala	Val 610	Thr	Lys	Glu	Glu	Tyr 615	Glu	Lys	Ala	Arg	Glu 620	Lys	Trp	Leu	Lys
Glu 625	ГЛЗ	Ala	Glu	Ser	Asn 630	Glu	Lys	Ala	Gln	Lys 635	Glu	Leu	Ala	Ser	His 640
Val	Lys														

- (2) INFORMATION FOR SEQ ID NO:49:
  - (i) SEQUENCE CHARACTERISTICS:
    - (A) LENGTH: 27 base pairs (B) TYPE: nucleic acid

    - (C) STRANDEDNESS: both (D) TOPOLOGY: linear
  - (ii) MOLECULE TYPE: cDNA
  - (iii) HYPOTHETICAL: NO
  - (iv) ANTI-SENSE: NO
  - (ix) FEATURE:
    - (A) NAME/KEY: CDS
    - (B) LOCATION: 1..1932
  - (xi) SEQUENCE DESCRIPTION: SEO ID NO:49:

#### GCCGGATCCG GWGTWCTTGC WGCWTGC

27

- (2) INFORMATION FOR SEQ ID NO:50:
  - (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 21 base pairs
    (B) TYPE: nucleic acid
    (C) STRANDEDNESS: both
    (D) TOPOLOGY: linear
  - (ii) MOLECULE TYPE: cDNA
  - (iii) HYPOTHETICAL: NO
  - (iv) ANTI-SENSE: NO
  - (ix) FEATURE:
    - (A) NAME/KEY: CDS (B) LOCATION: 1..1932
  - (xi) SEQUENCE DESCRIPTION: SEQ ID NO:50:

# TACAAGAGAC TACTTGGATC C

21 (2) INFORMATION FOR SEO ID NO:51:

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(ii) MOLECULE TYPE: cDNA
(iii) HYPOTHETICAL: NO
(iv) ANTI-SENSE: NO

(ix) FEATURE:
(A) NAME/KEY: CDS
(B) LOCATION: 1..1932

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(i)	SEQUENCE CHARACTERISTICS:  (A) LENGTH: 30 base pairs (B) TTPS: nucleic acid (C) STRANDEDINESS: both (D) TOPOLOGY: linear	
(ii)	MOLECULE TYPE: cDNA	
(iii)	HYPOTHETICAL: NO	
(iv)	ANTI-SENSE: NO	
(ix)	FEATURE: (A) NAME/KEY: CDS (B) LOCATION: 11932	
(xi)	SEQUENCE DESCRIPTION: SEQ ID NO:51:	
ACCGGATC	CT GCCAACAAGC CTAAATATTC	30
(2) INFO	RMATION FOR SEQ ID NO:52:	
(i)	SEQUENCE CHARACTERISTICS: (A) LENGTH: 30 base pairs (B) TTPS: nucleic acid (C) STRANDEDNESS: both (D) TOPOLOSY: linear	
(ii)	MOLECULE TYPE: cDNA	
(iii)	HYPOTHETICAL: NO	
(iv)	ANTI-SENSE: NO	
(ix)	PEATURE: (A) NAME/KEY: CDS (B) LOCATION: 11932	
(xi)	SEQUENCE DESCRIPTION: SEQ ID NO:52:	
TTTGGATC	CG TTGGTTTAGC AAAATCGCTT	30
(2) INFO	RMATION FOR SEQ ID NO:53:	
(i)	SEQUENCE CHARACTERISTICS:  (A) LENGTH: 18 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: both (D) TOPOLOGY: linear	

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(xi) SEQUENCE DESCRIPTION: SEQ ID NO:53:	
CTATACCTTG GTTCCTCG	. 1
(2) INFORMATION FOR SEQ ID NO:54:	
(i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 26 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: both (D) TOPOLOGY: linear	
(ii) MOLECULE TYPE: cDNA	
(iii) HYPOTHETICAL: NO	
(iv) ANTI-SENSE: NO	
(ix) FEATURE: (A) NAME/KEY: CDS (B) LOCATION: 11932	
(xi) SEQUENCE DESCRIPTION: SEQ ID NO:54:	
TTTGGATTCG GAATTTCACG AGTAGC	26
(2) INFORMATION FOR SEQ ID NO:55:	
(i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 1929 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: both (D) TOPOLOGY: linear	
(ii) MOLECULE TYPE: cDNA	
(iii) HYPOTHETICAL: NO	
(iv) ANTI-SENSE: NO	
(vi) ORIGINAL SOURCE:  (A) ORGANISM: Streptococcus pneumoniae (B) STRAIN: R6	
(vii) IMMEDIATE SOURCE: (B) CLONE: pad1 (poxB)	
(ix) FEATURE: (A) NAME/KEY: CDS (B) LOCATION: 1541929	
(xi) SEQUENCE DESCRIPTION: SEQ ID NO:55:	
CTGTATTAGA ATAGAGAATA GAGAGTTTTG AGCAGATTTT TAGAAAAGTC AGCATAAATA	60
TGATACAGTG GAATAGTAAA AATTTGGAGA ACGTTTCCAA TTCTATGTAA TCGTATTCTC	120
CAAGITTAAA AAAATTGAAG GAGAGITATC ATT ATG ACT CAA GGG AAA ATT ACT Met Thr Gln Gly Lys 11e Thr ${}^{1}{}_{1}$	174
GCA TCT GCA GCA ATG CTT AAC GTA TTG AAA ACA TGG GGC GTA GAT ACA	222

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Ala	Ser	Ala 10	Ala	Met	Leu	Asn	Val 15	Leu	Lys	Thr	Trp	Gly 20		Asp	Thr	
							Thr					Met			TTG Leu	270
	Glu														ACA Thr 55	318
	GCT Ala															366
	GCA Ala															414
	TAC Tyr															462
TCA Ser	CGT Arg 105	CCA Pro	GTT Val	AAC Asn	GAA Glu	TTG Leu 110	AAC Asn	ATG Met	GAT Asp	GCT Ala	TTC Phe 115	CAA Gln	GAG Glu	CTT Leu	AAC Asn	510
	AAC Asn															558
	GCT Ala															606
	TCT Ser														GGT Gly	654
	CAA Gln															702
	TCA Ser 185															750
	GAA Glu															798
	GGT Gly															846
AAA Lys	GCA Ala	CCA Pro	ATC Ile 235	ATC Ile	ACA Thr	ACT Thr	GGT Gly	AAA Lys 240	AAC Asn	TTT Phe	GAA Glu	GCT Ala	TTC Phe 245	GAA Glu	TGG Trp	894
	TAT Tyr															942
GCC Ala	AAC Asn	GAA Glu	GTG Val	GTC Val	TTT Phe	GAA Glu	GCA Ala	GAC Asp	ACA Thr	GTT Val	CTT Leu	TTC Phe	CTT Leu	GGT Gly	TCA Ser	990

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	265					270					275					
														GAA Glu	AAA Lys 295	1038
														CAT His 310		1086
														GCT Ala		1134
								Ser						GCA Ala		1182
														GAA Glu		1230
														ATC Ile		1278
AAA Lys	CAT His	GCT Ala	GAT Asp	CAA Gln 380	GAC Asp	GCT Ala	ATC	TAC Tyr	TCA Ser 385	CTC Leu	GAC Asp	GTC Val	GGT Gly	AGC Ser 390	ACT Thr	1326
ACT Thr	CAA Gln	ACA Thr	TCT Ser 395	ACT Thr	CGT Arg	CAC His	CTC Leu	CAC His 400	ATG Met	ACA Thr	CCT Pro	AAG Lys	AAT Asn 405	ATG Met	TGG Trp	1374
CGT Arg	ACA Thr	TCT Ser 410	CCG Pro	CTC	TTT Phe	GCG Ala	ACA Thr 415	ATG Met	GGT Gly	ATT Ile	GCC Ala	CTT Leu 420	CCT Pro	GGT Gly	GGT Gly	1422
ATC Ile	GCT Ala 425	GCT Ala	AAG Lys	AAA Lys	GAC Asp	ACT Thr 430	CCA Pro	GAT Asp	CGC Arg	CAA Gln	GTA Val 435	TGG Trp	AAC Asn	ATC Ile	ATG Met	1470
														AAC Asn		1518
CAA Gln	TAC Tyr	GAC Asp	CTT Leu	CCA Pro 460	GTT Val	ATC Ile	AAC Asn	CTT Leu	GTC Val 465	TTC Phe	TCA Ser	AAT Asn	GCT Ala	GAG Glu 470	TAC Tyr	1566
GGC Gly	TTC Phe	ATC Ile	AAG Lys 475	AAC Asn	AAA Lys	TAC Tyr	GAA Glu	GAT Asp 480	ACA Thr	AAC Asn	AAA Lys	CAC His	TTG Leu 485	TTT	GGT Gly	1614
GTT Val	GAC Asp	TTC Phe 490	ACA Thr	ATC Ile	GCT Ala	GAC Asp	TAC Tyr 495	GGT Gly	AAC Asn	CTT Leu	GCG Ala	GAA Glu 500	GCT Ala	CAC His	GGA Gly	1662
Ala	Val 505	Gly	Phe	Thr	Val	Asp 510	Arg	Ile	Asp	Asp	Ile 515	Asp	Ala	GTT Val	Val	1710
GCA Ala 520	GAT Asp	GCT Ala	GTT Val	AAA Lys	TTG Leu 525	AAC Asn	ACA Thr	GAT Asp	GGT Gly	AAA Lys 530	ACT Thr	GIT Val	GTC Val	ATC Ile	ĠAT Asp 535	1758

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1806 1854 1902

					His			Leu		Val					Leu
								GCT Ala 560							
			Glu					TTC Phe							
					GCA Ala										
(2)			SEQU (A (B	ENCE	NGTH PE:	RACT	ERIST	TICS ino a		5					
		;;) ;			POLOG										
						_		: SE(	Q ID	NO:	56:				
Met 1	Thr	Gln	Gly	Lys 5	Ile	Thr	Ala	Ser	Ala 10	Ala	Met	Leu	Asn	Val 15	Leu
Lys	Thr	Trp	Gly 20	Val	Asp	Thr	Ile	Tyr 25	Gly	Ile	Pro	Ser	Gly 30	Thr	Leu
Ser	Ser	Leu 35	Met	Asp	Ala	Leu	Ala 40	Glu	Asp	Lys	Asp	Ile 45	Arg	Phe	Leu
Gln	Val 50	Arg	His	Glu	Glu	Thr 55	Gly	Ala	Leu	Ala	Ala 60	Val	Met	Gln	Ala
Lys 65	Phe	Gly	Gly	Ser	lle 70	Gly	Val	Ala	Val	Gly 75	Ser	Gly	Gly	Pro	Gly 80
Ala	Thr	His	Leu	Ile 85	Asn	Gly	Val	Tyr	Asp 90	Ala	Ala	Met	Asp	Asn 95	Thr
Pro	Phe	Leu	Ala 100	Ile	Leu	Gly	Ser	Arg 105	Pro	Val	Asn	Glu	Leu 110	Asn	Met
Asp	Ala	Phe 115	Gln	Glu	Leu	Asn	Gln 120	Asn	Pro	Met	Tyr	Asn 125	Gly	Ile	Ala
Val	Tyr 130	Asn	Lys	Arg	Val	Ala 135	Tyr	Ala	Glu	Gln	Leu 140	Pro	Lys	Val	Ile
Asp 145	Glu	Ala	Cys	Arg	Ala 150	Ala	Ile	Ser	Lys	Lys 155	Gly	Pro	Ala	Val	Val 160
Glu	Ile	Pro	<b>V</b> al	Asn 165	Phe	Gly	Phe	Gln `	Glu 170	Ile	Asp	Glu	Asn	Ser 175	Tyr
Tyr	Gly	Ser	Gly 180	Ser	Tyr	Glu	Arg	Ser 185	Phe	Ile	Ala	Pro	Ala 190	Leu	Asn
Glu	Val	Glu	Ile	Asp	Lys	Ala	Val	Glu	Ile	Leu	Asn	Asn	Ala	Glu	Arq

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195 200 205 Pro Val Ile Tyr Ala Gly Phe Gly Gly Val Lys Ala Gly Glu Val Ile 210 215 220 Thr Glu Leu Ser Arg Lys Ile Lys Ala Pro Ile Ile Thr Thr Gly Lys Asn Phe Glu Ala Phe Glu Trp Asn Tyr Glu Gly Leu Thr Gly Ser Ala 245 250 255 Tyr Arg Val Gly Trp Lys Pro Ala Asn Glu Val Val Phe Glu Ala Asp 260 265 270 Thr Val Leu Phe Leu Gly Ser Asn Phe Ala Phe Ala Glu Val Tyr Glu 275 280 285 Ala Phe Lys Asn Thr Glu Lys Phe Ile Gln Val Asp Ile Asp Pro Tyr 290 295 300 Lys Leu Gly Lys Arg His Ala Leu Asp Ala Ser Ile Leu Gly Asp Ala 305 310 315 Gly Gln Ala Ala Lys Ala Ile Leu Asp Lys Val Asn Pro Val Glu Ser 325 330 335 Thr Pro Trp Arg Ala Asn Val Lys Asn Asn Gln Asn Trp Arg Asp 340 345 350 Tyr Met Asn Lys Leu Glu Gly Lys Thr Glu Gly Glu Leu Gln Leu Tyr 355 360 365 Gln Val Tyr Asn Ala Ile Asn Lys His Ala Asp Gln Asp Ala Ile Tyr 370 380 Ser Leu Asp Val Gly Ser Thr Thr Gln Thr Ser Thr Arg His Leu His 385 390 395 400 Met Thr Pro Lys Asn Met Trp Arg Thr Ser Pro Leu Phe Ala Thr Met
405 410 415 Gly Ile Ala Leu Pro Gly Gly Ile Ala Ala Lys Lys Asp Thr Pro Asp 420 425 430 Arg Gln Val Trp Asn Ile Met Gly Asp Gly Ala Phe Asn Met Cys Tyr
435 440 445 Pro Asp Val Ile Thr Asn Val Gln Tyr Asp Leu Pro Val Ile Asn Leu 450 . 455 460 Val Phe Ser Asn Ala Glu Tyr Gly Phe Ile Lys Asn Lys Tyr Glu Asp 465 470 480 Thr Asn Lys His Leu Phe Gly Val Asp Phe Thr Ile Ala Asp Tyr Gly Asn Leu Ala Glu Ala His Gly Ala Val Gly Phe Thr Val Asp Arg Ile Asp Asp Ile Asp Ala Val Val Ala Asp Ala Val Lys Leu Asn Thr Asp 515 520 525 Gly Lys Thr Val Val Ile Asp Ala Arg Ile Thr Gln His Arg Pro Leu 530 540

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Pro Val Glu Val Leu Asp Leu Val Pro Asn Leu His Ser Glu Glu Ala 545  $\phantom{\bigg|}550\phantom{\bigg|}$ 

Arg Leu Phe Leu Glu Glu Glu Glu Leu His Pro Arg Ala Ile Lys  $580 \hspace{1.5cm} 585 \hspace{1.5cm} 585 \hspace{1.5cm} 590 \hspace{1.5cm}$ 

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# WHAT IS CLAIMED IS:

- 1 1. A recombinant DNA molecule having the nucleotide sequence of SEQ ID
- 2 NO:46, or a hybridizable fragment thereof.
- 1 2. A recombinant DNA molecule having the nucleotide sequence of SEO ID
- 2 NO:5, or a hybridizable fragment thereof.
- 1 3. A recombinant DNA molecule having the nucleotide sequence of SEO ID
- 2 NO:7, or a hybridizable fragment thereof.
- 1 4. A recombinant DNA molecule having the nucleotide sequence of SEQ ID
- 2 NO:9, or a hybridizable fragment thereof.
- 1 5. A recombinant DNA molecule having the nucleotide sequence of SEQ ID
- 2 NO:11, or a hybridizable fragment thereof.
- 1 6. A recombinant DNA molecule having the nucleotide sequence of SEQ ID
- 2 NO:13, or a hybridizable fragment thereof.
- 1 7. A recombinant DNA molecule having the nucleotide sequence of SEO ID
- 2 NO:15, or a hybridizable fragment thereof.
- 1 8. A recombinant DNA molecule having the nucleotide sequence of SEQ ID
- 2 NO:17, or a hybridizable fragment thereof.
- 1 9. A recombinant DNA molecule having the nucleotide sequence of SEO ID
- 2 NO:19, or a hybridizable fragment thereof.
- 1 10. A recombinant DNA molecule having the nucleotide sequence of SEQ ID
- 2 NO:21, or a hybridizable fragment thereof.

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- 1 11. A recombinant DNA molecule having the nucleotide sequence of SEO ID
- 2 NO:55, or a hybridizable fragment thereof.
- 1 12. An isolated polypeptide having the amino acid sequence of SEO ID NO:47.
- 2 or an antigenic fragment thereof.
- 1 13. An isolated polypeptide having the amino acid sequence of SEQ ID NO:6, or
- 2 an antigenic fragment thereof.
- 1 14. An isolated polypeptide having the amino acid sequence of SEQ ID NO:8, or
- 2 an antigenic fragment thereof.
- 1 15. An isolated polypeptide having the amino acid sequence of SEO ID NO:10.
- 2 or an antigenic fragment thereof.
- 1. 16. An isolated polypeptide having the amino acid sequence of SEO ID NO:12.
- 2 or an antigenic fragment thereof.
- 1 17. An isolated polypeptide having the amino acid sequence of SEQ ID NO:14,
- 2 or an antigenic fragment thereof.
- 1 18. An isolated polypeptide having the amino acid sequence of SEQ ID NO:16,
- 2 or an antigenic fragment thereof.
- 1 19. An isolated polypeptide having the amino acid sequence of SEQ ID NO:18,
- 2 or an antigenic fragment thereof.
- 1 20. An isolated polypeptide having the amino acid sequence of SEQ ID NO:20.
- 2 or an antigenic fragment thereof.

- 21. An isolated polypeptide having the amino acid sequence of SEQ ID NO:22,
- 2 or an antigenic fragment thereof.
- 1 22. An isolated polypeptide having the amino acid sequence of SEQ ID NO:56.
- 2 or an antigenic fragment thereof.
- 1 23. A vaccine for protection of an animal subject from infection with a Gram
- 2 positive bacterium comprising a vector containing a gene encoding an exported
- 3 protein of a Gram positive bacterium operably associated with a promoter capable
- 4 of directing expression of the gene in the subject, in which the exported protein is
- 5 selected from the group consisting of an adhesion associated protein, a virulence
- 6 determinant, a toxin and an immunodominant protein.
- 1 24. A vaccine for protection of an animal subject from infection with a Gram
- 2 positive bacterium comprising a vector containing a gene encoding an exported
- 3 protein which is an antigen common to many strains of a species of Gram positive
- 4 bacterium operably associated with a promoter capable of directing expression of
- 5 the gene in the subject.
- 1 25. The vaccine of claim 23 or 24 in which the Gram positive bacterium is a S.
- 2 pneumoniae.
- 1 26. The vaccine of claim 23 or 24 in which the protein encoded by the gene is an
- 2 adhesin.
- 27. The vaccine of claim 25 in which the protein encoded by the gene is an
- 2 adhesin.
- 1 28. A vaccine for protection of an animal subject from infection with a S.
- 2 pneumoniae comprising a vector containing a gene encoding an exported protein of
- 3 a S. pneumoniae operably associated with a promoter capable of directing of

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- 4 directing expression of the gene in an animal subject, in which the gene contains a
- 5 nucleotide sequence selected from the group consisting of SEO ID NOS: 1, 3, 5,
- 6 7, 9, 11, 13, 15, 17, 19, 22, 46, 55, amiA and ponA.
- 1 29. The vaccine of claim 23, 24 or 28 in which the animal subject is a human.
- 1 30. A vaccine for protection of an animal subject from infection with a Gram
- 2 positive bacterium comprising an immunogenic amount of an exported protein of a
- 3 Gram positive bacterium and an adjuvant, in which the exported protein is selected
- 4 from the group consisting of an adhesion associated protein, a virulence
- 5 determinant, a toxin and an immunodominant protein,
- 1 31. A vaccine for protection of an animal subject from infection with a Gram
- 2 positive bacterium comprising an immunogenic amount of an exported protein
- 3 which is an antigen common to many strains of a species of Gram positive
- 4 bacterium and an adjuvant.
- 1 32. The vaccine of claim 30 or 31 in which the Gram positive bacterium is a S.
- 2 pneumoniae.
- 1 33. The vaccine of claim 30 or 31 in which the protein encoded by the gene is an
- 2 adhesin.
- 1 34. The vaccine of claim 32 in which the protein encoded by the gene is an
- 2 adhesin.
- 1 35. A vaccine for protection of an animal subject from infection with a S.
- 2 pneumoniae comprising an immunogenic amount of an exported protein of a S.
- 3 pneumoniae and an adjuvant, in which the exported protein contains an amino acid
- 4 sequence selected from the group consisting of SEQ ID NOS: 2, 4, 6, 8, 10, 12,
- 5 14, 16, 18, 20, 22, 47, 56, PonA and AmiA.

1 36. The vaccine of claim 30, 31, or 35 in which the animal subject is a human.

- 1 37. A method for identifying a portion of a gene encoding an adhesion associated
- exported protein of a Gram positive bacterium comprising the steps of:
- a. translationally inserting a DNA molecule obtained from a Gram positive
- 4 bacterium upstream of and in an open reading frame with an indicator
- 5 protein gene lacking its signal sequence and promoter in a vector in which
- 6 duplication mutagenesis of the Gram positive DNA molecule can occur,
- 7 wherein the indicator protein is non-functional unless exported by a
- 8 bacterium:
- b. introducing the vector into the Gram positive bacterium;
- 10 c. growing the Gram positive bacterium whereby a fusion protein of an
- 11 exported protein of the gram positive bacterium and the indicator protein
- 12 can be expressed:
- d. selecting bacteria in which the indicator protein is functional, indicating
- 14 export of the indicator protein;
- 15 e. screening for loss of adherence of the Gram positive bacterium to a
- 16 eukaryotic cell to which it normally adheres; and
- 17 f. selecting Gram positive bacteria that demonstrate loss of adherence:
- 18 whereby Gram positive bacteria containing a mutated gene encoding an exported
- 19 adhesion associated protein are selected.
  - 38. A method for identifying a portion of a gene encoding an exported protein
- 2 that is a virulence determinant of a Gram positive bacterium comprising the steps
- 3 of:
- 4 a. translationally inserting a DNA molecule obtained from a Gram positive
- 5 bacterium upstream of and in an open reading frame with an indicator
- 6 protein gene lacking its signal sequence and promoter in a vector in which
- 7 duplication mutagenesis of the Gram positive DNA molecule can occur,
- 8 wherein the indicator protein is non-functional unless exported by a
- 9 bacterium;

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- b. introducing the vector into the Gram positive bacterium;
- 11 c. growing the Gram positive bacterium whereby a fusion protein of an
- 12 exported protein of the gram positive bacterium and the indicator protein
- 13 can be expressed;
- d. selecting bacteria in which the indicator protein is functional, indicating
- 15 export of the indicator protein;
- 16 e. screening for loss of virulence of the Gram positive bacterium in an
- 17 animal LD<sub>50</sub> model; and
- 18 f. selecting Gram positive bacteria that demonstrate loss of virulence:
- 19 whereby Gram positive bacteria containing a mutated gene encoding an exported
- 20 protein virulence determinant are selected.
  - 39. The method according to claim 37 or 38 in which the indicator protein is
- 2. Escherichia coli PhoA.
- 1 40. The method according to claim 37 or 38 in which the Gram positive
- 2 bacterium is a S. pneumoniae.
- 1 41. The method according to claim 37 or 38 in which the exported protein is an
- 2 adhesin.
- 1 42. A vaccine for protection of an animal subject from infection with a Gram
- 2 positive bacterium comprising a vector containing a gene identified according to
- 3 the method of claim 37 or 38 operably associated with a promoter capable of
- 4 directing expression of the gene in an animal subject.
- 1 43. A vaccine for protection of an animal subject from infection with a Gram
- 2 positive bacterium comprising an immunogenic amount of a protein encoded by a
- 3 gene identified according to the method of claim 37 or 38 and an adjuvant.

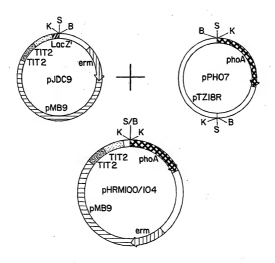
- 1 44. An antibody or fragment thereof reactive with a protein having an amino acid
- 2 sequence selected from the group consisting of SEQ ID NOS: 2, 6, 8, 10, 12, 14,
- 3 16, 18, 20, 22, 47 and 56,
- 1 45. A method for protecting a subject from infection with a Gram positive
- 2 bacterium comprising administering an immunogenic dose of a vaccine of claim
- 3 23, 24, 28, 30 or 31.
- 1 46. A method for protecting a subject from infection with a Gram positive
- 2 bacterium comprising administering an immunogenic dose of a vaccine of claim
- 3 42.
- 1 47. A method for protecting a subject from infection with a Gram positive
- 2 bacterium comprising administering an immunogenic dose of a vaccine of claim
- 3 43.
- 1 48. A method for diagnosing an infection with a Gram positive bacterium
- 2 comprising detecting the presence of a Gram positive bacterium with an antibody
- 3 or fragment thereof of claim 44.
- 1 49. A method for diagnosing an infection with a Gram positive bacterium
- 2 comprising detecting the presence of a Gram positive bacterium in a sample from
- a subject with a nucleic acid probe which is a hybridizable fragment of a
- 4 recombinant DNA molecule having a nucleotide sequence selected from the group
- 5 consisting of SEO ID NOS: 1, 5, 7, 9, 11, 13, 15, 17, 19, 21, 46 and 55.
- 1 50. A method for diagnosing an infection with a Gram positive bacterium
- 2 comprising detecting the presence of a Gram positive bacterium by polymerase
- 3 chain reaction using a primer which is a hybridizable fragment of a recombinant
- 4 DNA molecule having a nucleotide sequence selected from the group consisting of
- 5 SEQ ID NOS: 1, 5, 7, 9, 11, 13, 15, 17, 19, 21, 46 and 55.

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- 1 51. A method for protecting a subject from infection with a Gram positive
- 2 bacterium comprising administering a therapeutically effective dose of an antibody
- 3 of claim 43.
- 1 52. A method for protecting a subject from infection with a Gram positive
- 2 bacterium comprising administering a therapeutically effective dose of a Gram
- 3 positive adhesin encoded by the gene isolated according to claim 37.

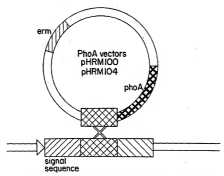
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FIG. IA

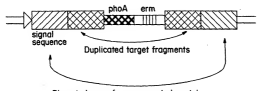


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FIG. 1B



Gene for an exported protein from S. pneumoniae



Disrupted gene for an exported protein

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FIG. 2

1 2 3 4

106.0 - 106.0

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FIG. 3

1 2 3

FIG.

120 120 120 9 9 9 9 8 OSLEATLGAD VTOEAATEAP GOTGASKILR NLFVPPTFVO ADGRNFGDMV KEKLVTYGDE WKDVNLADSO IYNTDEYVAY PDDELQVAST IVDVSNGKVI AQLGARHQSS YYODEADLPA IYTVVOEOER MLGRKIAVEV EOLRHYILAE FDGIIEFKAL SKDNLLQIVE RLRVTVKDAD KVGNAEQWKA EGAMGLVMKG DIDYTGDLEH PKVTQEDVDY LLGIVNNRFP YLSKEKTRED TASEFELGTP LSQEXLDHHK POKPSDIQAL ALLEILDPIR EGAASTLDYL RSQEVGLKII OSRRCKTTAS EFELGTLRKN IGLVLOEPFL YHGTIKSNIA MYOEISDEOV LILDEADEML NMGFLEDIEA LRDFKNGNLD VLVATDVAAR YKD 173 YYI 115 OGVOALYGPK ADILKSDIOD ILDSGEIIPE TLPSOMTEVO ONTVHFKD 108 TKVORVOSMK KASLSEESYR RPISKELIHY FDNLIATVES KELTTELVDO FAD 163 LSDNVGWGPD LDMPVDQTAT NYEKPSOEVL ASGCGWPSFS NFFSPEFMIR KALKLODIET KAPEHVRIAA GNNGTIRDES VDELTRGLKI RGFRAEGIHG DLDQNKRLRV FSTGOR 96 GYD 93 QAEGVTFPIH AENAAGEDWD PADKPLIDAA EPLFFAKOKF SVGFGAAREG RINSVLGELG SSNNIRLDVT DKVKEXLVDL VKVDDGSOAV NIINLLGGRV NIVDVDACMT GDGRMVFVLP RENKTYFGTT DSPVSERGSS KSGAHIVVGT PGRLLDLIKR TLLFSATMPD AIKRIGVOFM SSWAGLRPLI AGNSASDYNG SGDNPVTVSS IAQKAGFADY H 71 STSEKHLD 139 DRGROYRIGI AEFAKAKSAL KDEVNNITYF DQEAQKHLWD PSGYCHIDVT EGIYVDITTG SFIQELPOGY YAYDIPOD 78 NVSFGINOAV ETNRDWG 77 DRIAYASOLN DGLYNPEKAK TIGMDVYTNV DPLSINOOGN LMLADVNKRL SOPVSFDTGL 1 RLSWVTPGFS NVIDIOOLO DYHODYLRKN FINAYDOTFE SNAGTGKTEA VESAVSKLES OAAAAFVDAD IISRVPENRO AFVFGRTKRR GLDISGVTHV ESNITIDDIE SSIEKOIKAL 121 -121 н -4 121 -5 Exp9 a Exp9 b Exp6 Exp8 Exp1 Exp5 Exp7 Exp2 Exp3 Exp4

	4 ID	#SIM
EXPI I DRTA:ASQLNOQTGASKILRNLFVPPTFVQADGANFFGDNYKEKLVTKGOSMKDVNLADSQDGLVNPE:AKAEFAKAKSALQAEGV 85 DR+AX++O+NG GA+ +RNLFV P FV A K FGD+V L +YGDEW VNLAD-GDGL+N++KAKAEF KAK AL+A+GV	9	42
Amia 148 drsaysaqingkdgaalavralevrpoeysagektegduvaaqerygdemgayladgqoglenadkaeerkakkaleadgv 411	:	<b>!</b>
Expl 86 TFPHLDMPUDGTATTKVQRVQSLEATLGADWITDTQQLQKDEVNVITTRABAAGENAVGEDOFAD 163		
FILLLOFFULY** * KAJA NOSTER LG ***********************************		
FIG. 5B		
Exp2 1 TICHDUYTINUDGEACKHEMDIYNIDE:VAYPDDSELQVASTIVDVSNGKVIAQLCARHQSSNVSFGINQAVETINROMG 77	QT.	E
TTGHOYTINTOGEACKHIMDIYNTOGEYAXY DOBELQYASTIYOVSNGKY IAQLGARHQSSNYSTGINQAVETNROMG Pola 353 ttghoytintogeackhilmdiyntogeyaxy edbeelqyastiyovsngky laqlgarhqssnystgingavetnromg 369	100	100
FIG. 5C		
Exd3 1 DPLSINQGGNDRGRQYRTGIYYQDEADLPAIYTVVQEQERANLGRKIAVEVEQLRHYILAEDYHQDYLRKNPSGYCHIDVTDADKPL 86	. QI	2 9 Wis
DP 5-M GGND G GYR-G-YY D A+ + I + + +++ + VE E L+++ A5-VRGOYL KNP-GYCHID+ AD PL PHB 274 OPTSLMYGGNYTGQYRSGVYTDPAEXALKAEGGNYGLPLWERFERLKNFYDAEZFKGDYLIKNPNGYGHIDIRAADEPL 359	Ç	64
Exp3 87 idranterpsqevlkaslesestryt 112		
PilB 360 pcktyca-pocalarogrikantersnaperaalfed 397	E.	ğ
Exp3 113 QEAATEAPETRALOCTESSOLYPOTTOSPALFAXOCAAPESSAP1 163	;	;
Q++ATS +*** IV+ F GIVON +GGZEF + DK SCCGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGG	·	883
FIG. 5D		į
Exp4 25 viceignfespeernredoilefkalskbullqivelmladvukrissnniridovtokvkekjvologo 93	Q1 <b>*</b>	WT S.
V EL 4F PEFARR D 41 F+ L+X ++ 1 ++ML 4-Y L-X-TELLAVERISLAVERSENDRIVDEGYN 858	#	. 14

•	S.				WIS#	78	7/	#SIM	€	Ì			į		<b>5</b>	*STH	6
919	1				07.	89		ÇI.	36	:	3	9	:	3	7	619	ž
4 DOSCONVIINLLOGRYRIUDVDACATELAVTYKDADKYGNARGAWASGAAGUYAKGGOYQALYGRYADILKSDIQDILDSGZII 88	. G + 1 G + HI - PACTHIMATOND K + 1. K GA G1. G - CALEGRAD LKNGDI. + 189 ENGDLFWEILQANGDDEDIKELDACITELKUTVNDQKV DKDRKQLGASGVLEVGANGALGAIFGPRSSGLKTQMQDILAGRSPR 274	89 PETL2SQWTEWQQ iol BF S FV O	TSYG		12 DCRAVFVLPRENKTYFGTTDTDTTGGLEHPKVTQEDVDYLLGIVNNRFPESNITIDDISSSMAGLRPLI 80	DGRAVF +PRE KTY GTTDT Y LEHP++T ED DY++ +N FPE NIT +DIESSWAGLRPLI 278 DGRAVFAIPREGKTYVGTTDTVYKEALEHPRHTTEDRDVIKSINVHFPELNITANDIESSWAGLRPLI 346		1) ALLEIL DPVREGAAETLDYLRSOEVGLKIISGVNPVTVSSI 71	•	*	ZGEISDE	+-ER+ +G+VIQEF L++ +++ NIA- 542 ASLARQLGVVLQESTLERRS/RDNIALTRPGASMH EVVAAARLAGAHESTGQLFZGYDYMLGENGYGLSGGGR. 614		6 QIKALKSGAHIVVGTPGALLDLIKRKALKLQDIETLILDEADEMLANGFIEDIEALISRVPENRÇTLIFSATMPDAIKALGVQFHK 91	QALG GIVVGTGRALD +KK -L L + L-ALDKADEL MG-ELL	1 KEVFORTKRRPUTELINGERREGERACIEDINGER DEN WASHENDEN DEN VERTER DES NEW THEVEN THE TRANSPORTED TO	A +F RTK E++ +L G+ +++GD++Q R + L K+G LD+LATDVARGLD+ ++ V NYDLP D 265 ALIEVATKNATLEVARALDVEARGLOVERISLAMONALNGOMYOALRSCOTLERLEDGEALDILATDVARGLOVERISLAMONALNGORNOALRSCOTLERLEDGEALDILATDVARGLOVERISLAMONALNGORNOALRSCOTLERLEDGEALDILATDVARGLOVERISLAMONALNGORNO
F1G. 5E	PtsG	Exp5	PtsG	F1G. 5F	Exp6	GlpD	F1G 5G	) .	MgtB	FIG. 5H	Exp8	CyaB	FIG. 5I	Exp9	DeaD	P. 42	DeaD

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FIG. 6

1 2

106.0 ----

80.0



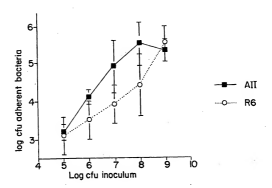
49.5 ---

32.5 ---

27.5 ---

18.5

FIG. 7



WO 95/06732



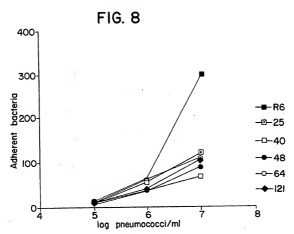
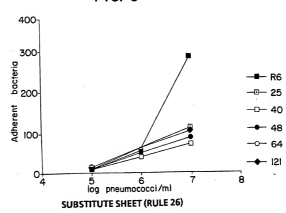


FIG. 9



F16.10

ProAspAlaLeuArgLysProThrCysSerLeuHisArgGluPheAspArgLeuSerGln AGGAGCCATAACGATTTTGAAGGCGTTGAATCGTATCATGAATACACAATACGACAA ArgSerHisAsnAspPheGluGlyValGluSerTyrHisGluTyrThrIleArgGln **ACTCTCACCATATCCTCATATGTAACAGAAGGATTACCAATTTTTAGGATTATCAATAATT** ThrLeuThrIleSerSerTyrValThrGluGlyLeuProIlePheArgLeuSerIleIle GTCCCTGTAAAGAGTCTATGGTCTGTGGAATGTAGGAAATTTTCTTACGTAGTTTACCAT ValProValLysSerLeuTrpSerValGluCysArgLysPheSerTyrValValTyrHis CCAGATGCGCTTCGAAAGCCAACGTGTTCTCTCCACCGAGAATTTGACCGCCTCTCTCAA AGTCAAGATC SerGlnAsp FIG.11A

gatgtacttgcagcatgctctggatcaggttcaagcgctaaaggtgagaagacattctca GlvValLeuAlaAlaCysSerGlySerGlySerSerAlaLysGlyGluLysThrPheSer tacatttatgagacagaccctgataacctcaactatttgacaactgctaaggctgcgaca PyrIleTyrGluThrAspProAspAsnLeuAsnTyrLeuThrThrAlaLysAlaAlaThr

gcaaatattaccagtaacgtggttgatggtttgctagaaaatgatcgctacgggaacttt AlaAsnIleThrSerAsnValValAspGlyLeuLeuGluAsnAspArgTyrGlyAsnPhe

ValProSerMetAlaGluAspTrpSerValSerLysAspGlyLeuThrTyrThrTyrThr stccgtaaggatgcaaaatggtatacttctgaaggtgaagaatacgcggcagtcaaagct

IleArgLysAspAlaLysTrpTyrThrSerGluGlyGluGluTyrAlaAlaValLysAla caagactttgtaacaggactaaaatatgctgctgataaaaaatcagatgctctttaccct gttcaagaatcaatcaaagggttggatgcctatgtaaaaggggaaatcaaagatttctca ValGlnGluSerIleLysGlyLeuAspAlaTyrValLysGlyGluIleLysAspPheSer

GlnaspPhevalThrGlyLeuLysTyralaalaaspLysSeraspalaLeuTyrPro

caagtaggaattaaggctctggatgaacagacagttcagtacactttgaacaaaccagaa GlnValGlyIleLysAlaLeuAspGluGlnThrValGlnTyrThrLeuAsnLySProGlu

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cottatttgttgaaatccattgtgaccaaatcctctgttgaatttgcgaaaatccgaac ProfyrLeuLeuLysSerIleValThrLysSerSerValGluPheAlaLysAsnProAsn

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yataccagcaaacctgcagaaaactttaaagatggtagccttacagcagctcgtctctat ASDThrSerLysProAlaGluAsnPheLysAspGlySerLeuThrAlaAlaArgLeuTyr

ccaacaagtgcaagtttcgcagagcttgagaagagtatgaaggacaatattgtctatact

ProThrSerAlaSerPheAlaGluLeuGluLysSerMetLysAspAsnIleValTyrThr caacaagactctattacgtatctagtcggtacaaatattgaccgtcagtcctataaatac 31nGlnAspSerIleThrTyrLeuValGlyThrAsnIleAspArgGlnSerTyrLysTyr 870

acatctaagaccagcgatgaacaaaaggcatcgactaaaaaggctctcttaaacaaggat ThrSerLysThrSerAspGluGlnLysAlaSerThrLysLysAlaLeuLeuAsnLysAsp

tecgteaggetattgeetttggttttgategtacageetatgeetedeagttgaatgga PheArgGlnAlaIleAlaPheGlyPheAspArgThrAlaTyrAlaSerGlnLeuAsnGly caaactggagcaagtaaaatcttgcgtaatctctttgtgcccaccaacatttgtccaagca 31nThrGlyAlaSerLysIleLeuArgAsnLeuPheValProProThrPheValGlnAla gatggtaaaaactttggcgatatggtcaaagagaaattggtcacttatggggatgaatgg AspGlyLysAsnPheGlyAspMetValLysGluLysLeuValThrTyrGlyAspGluTrp

aaggatgttaatcttgcagattctcaggatggtctttacaatccagaaaaagccaaggct LysAspValAsnLeuAlaAspSerGlnAspGlyLeuTyrAsnProGluLysAlaLysAla

## F16.110

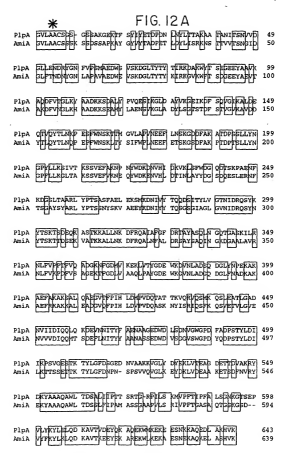
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                                                                                                                                    gatatgccagttgaccagacagcaactacaaagttcagcgcgtccaatctatgaaacaa
                                                                                                                                                                               AspMetProvalAspGlnThrAlaThrThrLysValGlnArgValGlnSerMetLysGln
                                                                                                                                                                                                                                                                        tccttggaagcaactttaggagctgataatgtcattattgatattcaacaactacaaaaa
                                                                                                                                                                                                                                                                                                              SerLeuglualaThrLeuglyalaAspAsnValIleIleAspIleglnGlnLeuglnLvs
                                                                                                                                                                                                                                                                                                                                                                                                                  gacgaagtaaacaatattacatattttgctgaaaatgctgctggggaagactgggattta
                                                                                                                                                                                                                                                                                                                                                                                                                                                       AspGluValAsnAsnIleThrTyrPheAlaGluAsnAlaAlaGlyGluAspTrpAspLeu
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           {	t Ser} {	t AspAsnValGlyTrpGlyProAspPheAlaAspProSerThrTyrLeuAspIleIle}
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          aaaccatctgtaggagaaagtactaaaacatatttagggtttgactcaggggaagataat
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     LysProSerValGlyGluSerThrLysThrTyrLeuGlyPheAspSerGlyGluAspAsn
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    gtagctgctaaaaaagtaggtctatatgactacgaaaaattggttactgaggctggtgat
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         ValalalalysLysValGlyLeuTyraspTyrGluLysLeuValThrGluAlaGlyAsp
                                                                                                                                                                                                                                                                                                                                                                  1410
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       1470
                                                                                                                                                                                                                              1350
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           1450
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tocttttttacatttgtaaagaaagattctaaatgtact

# F16.11D

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gagactacagatgttgctaaacgctatgataaatacgctgcagcccaagcttggttgaca
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       GluLysTrpMetLysGluLysGluGluSerAsnLysLysAlaGluAspLeuAlaLys
                                            GluThrThrAspValAlaLySArgTyrAspLySTyrAlaAlaAlaGlnAlaTrpLeuThr
                                                                                                                                         gatagtgctttgattattccaactacatctcgtacagggcgtccaatcttgtctaagatg
                                                                                                                                                                                     AspSerAlaLeullelleProThrThrSerArgThrGlyArgProIleLeuSerLysMet
                                                                                                                                                                                                                                                                                        gtaccatttacaataccatttgcattgtcaggaaataaaggtacaagtgaaccagtcttg
                                                                                                                                                                                                                                                                                                                               Val ProPheThrIleProPheAlaLeuSerGlyAsnLysGlyThrSerGluProValLeu
                                                                                                                                                                                                                                                                                                                                                                                                                            tataaatacttggaacttcaagacaaggcagtcactgtagatgaataccaaaaagctcag
                                                                                                                                                                                                                                                                                                                                                                                                                                                                         TyrLysTyrLeuGluLeuGlnAspLysAlaValThrValAspGluTyrGlnLysAlaGln
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                gaaaatggatgaaagaaagaagaagagtctaataaaaaggctcaagaagatctcqcaaaa
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          catgtgaaataactgttgcaaaatataaagaaaggatttagtatttctcttgaatgctgaa
                                                                                                                                                                                                                                       1790
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          1890
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  2010
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      HisValLysEnd
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       1990
```



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### FIG. 12B

plpA		OH BH	rio Pa	684	No.	643
AmiA		Spit Fet	H-1	NO9		659
SpoOKA		TOTAL SOL	1800	<del>224</del>	6/1	545
HbpA			304			547
DciAE		2/02) E/A	box	-	(As	543
OppA(Ec)						543
TraC		00	-	251		543
DopA			DOM:	·	<b></b>	<b>53</b> 5
PrgZ				Cont.	-	545
OppA (St)		· · · · · · · · · · · · · · · · · · ·		-	-	542
SarA		78/0 BN	10.01			624
	O 50	00 20 150		350 450 250 450	00 600 7 550 650	™ 700

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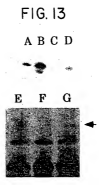


FIG. 14

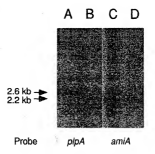
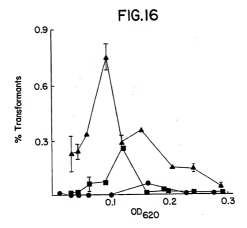
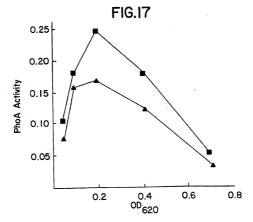


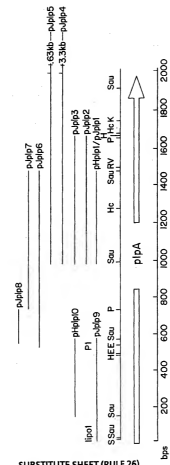
FIG. 15

Chromosomal gene construction	Strain	Percent of control
plpA	R6x	100.0 ± 17.6
plpA phoA E	SPRU98	12.5±3.2
plpA E	SPRUIO7	6.3±1.6
plpA phoA E	SPRU58	49.3±0.8
plpA E	SPRU 122	7.8±1.6
amiA amiC	R6x	100.0±28.2
amiA phoA E	SPRUI2I	116.3±16.1
amiA E	SPRUII4	130.2± 16.1
amiA amiC E	SPRUI48	371.5±31.9





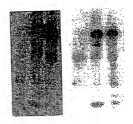
F16.18



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PCT/US94/09942 WO 95/06732

> 23/29 FIG. 19A ABC DEF



### Membrane Cytoplasm

A, D R6, parent B, E Pad1 C, F Pad1b

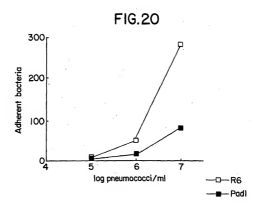
FIG. 19B

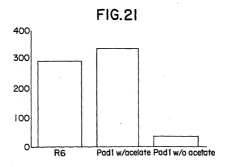
ABC



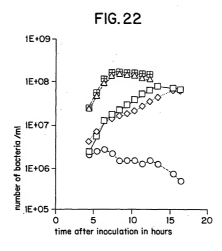
Membrane A, R6, parent B, Pad1 C, Pad1b

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F16. 23A

CTGTATTAGAATAGAATAGAGAGTTTTGAGCAGAŢTTTTAGAAAAGTCAGCATAAATATGATACAGTG 20

GAATAGTAAAAATTTGGAGAACGTTTCCAATTCTATGTAATCGTATTCTCAATTCTAAAAAAATTGAAG

**GAGAGTT ATCATTATGACTCAAGGGAAAATTACTGCATCTGCAATGCTTAAACGTÀTTGAAAACATG** 190

П > z μ, Σ ഗ K Н × Ö 0

GGCGTAGATACAATCTACGGTATCCCATCAGGAACACTCAGCTCATTGATGGACGCTTTGGCTGAAGACA 270 Д Σ ß S 250 Ö S Д 230 Ü

AAGATATCCGCTTCTTACAAGTTCGCCACGAAGAGACAGGTGCTCTTGCAGCGGTTATGCAAGCTAAATT 330 310 ы Ħ ø

CGCGGCTCAATCGGGGTTGCAGTTGGTTCAGGTGGTCCAGGTGCGACTCACTTGATTAACGGTGTTTAC 410 Ö 390 ρ. Ö Ö ď 370 Ö Ö

GATGCAGCTATGGATAACACTCCATTCCTAGCGATCCTTGGATCACGTCCAGTTAACGAATTGAACATGG 470 ρ, ĸ ഗ Ö ᆸ 450 ſ. Д 430

ATGCTTTCCAAGAGCTTAACCAAAACCCAATGTACAACGGTATCGCTGTTTACAACAAAAAGGTGTAA 550 530 O z z 510 z ы FIG. 23B

610 590

CGCTGAGCAATTGCCAAAAGTAATTGACGAAGCCTGCCGTGCTACAATTTCTAAAAAAGGTCCAGCTGTT Ø æ æ ш >

GTTGAAATTTCCAGTAAACTTCGGTTTCCAAGAAATCGACGAAAACTCATACGGTTCAGGTTCAATACG 670

ш ίĊ G S G > > S z ы Ω ы o 124 Ü 14 z > ρ, н Ш

AACGCTCATTCATCGCTCCTGCTTTGAACGAAGTTGAAATCGACAAAGCTGTTGAAATCTTGAACAATGC 750 730 z ч 710

TGAACGCCCAGTTATCTATGCTGGATTTGGTGGTGTTAAAGCTGGTGAAGTGATTACTGAATTGTCACGT > U 810 × > Ö Ĺ, Ü K 790

910 AAAATCAAAGCACCAATCATCACAACTGGTAAAACTTTGAAGCTTTCGAATGGAACTATGAAGGTTTGA > z 890 Ĺ K ы z 870 v Н н 850

CAGGTTCTGCTTACCGTGTTGGATAGCAAACCAGCCAACGAAGTGGTCTTTGAAGCAGACACAGAGTTCTTTT 970 ш > 950 Ø v 930 >

1050 CCTTGGTTCAAACTTCGCATTTGCTGAAGTTTACGAAGCATTCAAGAACACTGAAAAATTCATACAAGTC ø ы 1030 Н z × × 1010 ш . . 066

**GGTGATGGAGCATTCAACATGTGCTACCCAGACGTTATCACAAACGTTCAATACGACCTTCCAGTTATCA** 

×

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F16, 23C

SATATCGACCCTTACAAACTTGGTAAACGTCATGCCCTTGACGCTTCAATCCTTGGTGATGCTGGTCAAG

1090

1070

1110

Ö

ч

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1330 CAGCTAAAGCTATCCTTGACAAAGTAAACCCAGTTGAATCAACTCCATGGTGGCGTGCAAACGTTAAGAA CAACCAAAACTGGCGTGATTACATGAACAAACTCGAAGGTAAAACTGAGGGTGAATTGCAATTGTATCAA AAACATCTACTCGTCACCTCCACATGACACCT**AAGAAT**ATGTGGCGTACATCTCCGCTCTTTGCGACAAT z o 1530 1390 1250 ч Д 1450 E E ĸ 1170 1310 , 3 Д Д H × 1510 1230 1370 Ŋ Ω v ď ы z ш Д > ... ø 1150 A Z 1290 1430 4 × œ ď, z > Σ Σ H × 1210 × 1350 1490 Ω ĸ 1270 1130 z

Ą

F16. 23D

z Ω 1590 ы z ſı, 1570 O × K, z

TGGTGTTGACTTCACAATCGCTGACTACGGTAACCTTGCGGAAGCTCACGGAGCTGTTGGATTCACAGTT 1670 Ö ы 1650 L A Ü Ω K 1630

GACCGTATCGACGACATCGATGCAGTTGTTGCAGATGCTGTTAAATTGAACAAAGAAGGTAAAACTGTTG Z 1730 K Δ A > 1710 1690

TCATCGATGCTCGCATCACTCAACACCGTCCACTTCCAGTAGAAGTACTTGAATTGGATCCAAAACTTCA 1810 ы > 1790 > 4 Д ĸ × 0 1770 H

CTCAGAAGAAGCTATCAAAGCCTTCAAGGAAAAATACGAAGCAGAAGAACTCGTACCATTCCGTCTCTTC 1870 ы 1850 1830

1950 1930 Þ S 1910 O ч G

2045 GAGGAGAGTAACATG